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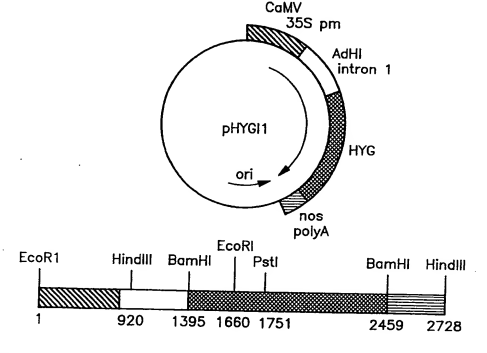
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(54) Title: FERTILE TRANSGENIC CORN PLANTS



(57) Abstract

Fertile transgenic Zea mays (corn) plants which stably express recombinant DNA which is heritable are provided wherein said DNA preferably comprises a recombinant gene which encodes a seed storage protein, so that the amino acid profile of the corn is improved.

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FERTILE TRANSGENIC CORN PLANTS

Field of the Invention

This invention relates to fertile transgenic

5 plants of the species Zea mays (oftentimes referred to herein as maize or corn). The invention further relates to producing fertile transgenic plants via particle bombardment and subsequent selection techniques.

10 <u>Background of the Invention</u>

Genetic engineering of plants, which entails the isolation and manipulation of genetic material (usually in the form of DNA or RNA) and the subsequent introduction of that genetic material into a plant or plant cells, offers considerable promise to modern agriculture and plant breeding. Increased crop food values, higher yields, feed value, reduced production costs, pest resistance, stress tolerance, drought resistance, the production of pharmaceuticals, chemicals 20 and biological molecules as well as other beneficial traits are all potentially achievable through genetic engineering techniques. Once a gene has been identified, cloned, and engineered, it is still necessary to introduce it into a plant of interest in such a 25 manner that the resulting plant is both fertile and capable of passing the gene on to its progeny.

A variety of methods have been developed and are currently available for the transformation of various plants and plant cells with DNA. Generally, 30 these plants have been dicotyledonous, and some success has been reported with certain of the monocotyledonous cereals. However, some species have heretofore proven untransformable by any method. Thus, previous to this discovery, no technology had been developed which would permit the production of stably transformed Zea mays plants in which the introduced recombinant DNA is transmitted through at least one complete sexual cycle. This failure in the art is well documented in the literature and has been discussed in a number of recent

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reviews (I. Potrykus, <u>Trends in Biotechnology</u>, <u>7</u>, 269 (1989); K. Weising et al., <u>Ann. Rev. of Genetics</u>, <u>22</u>, 421 (1988); F. Cocking et al., <u>Science</u>, <u>236</u>, 1259 (1987)).

Some of the techniques attempted, or proposed, for introducing DNA into corn cells include electroporation, microinjection, microprojectile bombardment, liposome fusion, Agrobacterium-mediated transfer, macroinjection, and exposure to naked DNA in solution.

Manipulation of Ovule Tissue, G. Chapman et al., eds.,
Longman, Inc., New York (1985) at pages 197-269 and Y.
Ohta, PNAS USA, 83, 715 (1986) reported the introduction
of DNA into maize by mixing pollen grains with DNA
solutions, and applying the pollen to maize silks. In
these papers, there is no molecular data confirming the
introduction of the exogenous DNA into the corn cells.
Arntzen et al. in published European Patent Application
No. 275,069 describe the incubation of DNA with maize
pollen followed by pollination of maize ears and
formation of seeds. The plants derived from these seeds
were reported to contain the introduced DNA, but there
is no suggestion that the introduced DNA was transmitted
through a complete sexual cycle.

A. Graves et al., <u>Plant Mol. Biol.</u>, <u>3</u>, 43

(1986) reported <u>Agrobacterium</u>-mediated transformation of <u>Zea mays</u> seedlings. The evidence was based upon assays which can sometimes be unreliable. To date, there have been no further reported successes with pollen and Agrobacterium-mediated transfer techniques.

Microprojectile bombardment has been reported to yield transformed corn cells. The technique is disclosed in Sanford et al., Part. Sci. & Techn., 5, 27 (1987) as well as in published European patent application number 331,855 of J.C. Sanford et al. which is based upon U.S. Serial No. 07/161,807, filed February

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29, 1988. Klein et al., Plant Physiol., 91, 440 (1989) describe production of transformed corn cells, using microprojectile bombardment. However, the cells used were not capable of regeneration into plants. 5 protocols have been published describing the introduction of DNA by a bombardment technique into cultures of regenerable maize cells of any type. No stable introduction of a gene has been reported that results from bombardment of maize callus followed by regeneration fertile plants and transmission of the introduced gene through at least one sexual cycle. D. McCabe et al., in published European patent application No. 270,356, disclose the bombardment of maize pollen with DNA, the application of the pollen to 15 silks, and the formation of seeds which reportedly contain the exogenous DNA. However, there is no evidence that the DNA was transmitted through a complete sexual cycle, and no further results have been reported by this group.

20 Electroporation of corn protoplasts has been reported to result in transformed cells by M.E. Fromm et al., Nature, 319, 791 (1986) although these cells did not provide regenerated plants. Electroporation of corn protoplasts was also reported by C. Rhodes et al., 25 Science, 240, 204 (1988). Although the recipient cells were transformed and, in the latter case, were able to regenerate into plants, the plants themselves were sterile. In addition, methods for the production of the cell line used by Rhodes et al. were not reproducible.

A further stumbling block to the successful production of fertile transgenic maize plants has been in selecting those few transformants in such a manner that neither the regeneration capacity (in the case of protoplasts or cell cultures) nor the fertility of the 35 transformants are destroyed. Due to the generally low level of transformants produced by a transformation technique, some sort of selection procedure is often

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necessary. However, selection generally entails the use of some toxic agent, e.g., a herbicide or antibiotic, which may be detrimental to either the regenerability or the resultant plant fertility.

On the other hand, it has been known that untransformed corn protoplasts, cultured cells and callus at least can be regenerated to form mature plants and that the resulting plants are often fertile. For example, R.D. Shillito et al., Bio/Technology, 7, 581 10 (1989), and L.M. Prioli et al., Bio/Technology, 7, 589 (1989) discuss methods for producing protoplasts from cell cultures and recovering fertile plants therefrom. C.A. Rhodes et al., Bio/Technology, 6, 56 (1988) disclose attempts to regenerate maize plants from protoplasts isolated from embryogenic maize cell cultures.

However, it has not been possible for the art worker to determine which maize tissues or cultures are appropriate recipients for exogenous DNA, e.g., contain 20 a useful number of cells which are receptive to and which will stably integrate the exogenous DNA, and at the same time be a part of the germline, i.e., part of the cell lineage that leads to the next plant generation.

The art is thus faced with a dilemma. certain transformation techniques have been proposed or reported to produce transformed maize cells and certain cells and tissues have been proposed to be potential recipients due to their ability to regenerate plants, the art has failed to find a combination of techniques which would successfully produce transformed maize plants able to transmit the introduced DNA through one complete sexual cycle.

It is thus an object of the present invention 35 to produce fertile, stably-transgenic, Zea mays plants and seeds which transmit the introduced gene to progeny. It is a further object to produce such stably-transgenic 15

plants and seeds by a particle bombardment and a selection process which results in a high level of viability for at least a portion of the transformed cells. It is a further object to produce fertile, stably-transgenic plants of other graminaceous cereals besides maize.

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Summary of the Invention

The present invention relates to fertile transgenic Zea mays plants containing recombinant DNA, preferably chromosomally integrated recombinant DNA, which is heritable by progeny thereof.

The invention further relates to all products 20 derived from transgenic Zea mays plants, plant cells, plant parts, and seeds.

The invention further relates to transgenic Zea mays seeds stably containing recombinant DNA and progeny which have inherited the recombinant DNA. The invention further relates to the breeding of transgenic plants and the subsequent incorporation of recombinant DNA into any Zea mays plant or line.

The invention further relates to a process for producing fertile transgenic Zea mays plants containing recombinant DNA. The process is based upon microprojectile bombardment, selection, plant regeneration, and conventional backcrossing techniques.

The invention further relates to a process for producing fertile transformed plants of graminaceous

35 species other than Zea mays which have not been reliably transformed by traditional methods such as electroporation, Agrobacterium, injection, and previous

ballistic techniques.

The invention further relates to regenerated fertile mature maize plants obtained from transformed embryogenic tissue, transgenic seeds produced therefrom, and R1 and subsequent generations.

A preferred embodiment of the present invention is a fertile, transgenic corn plant that has been stably transformed with a recombinant, chimeric gene which is expressed as a seed storage protein, e.g., as the 10 kD zein protein, so that the level of at least one amino acid is elevated above that present in the parent, nontransformed lines. Expression of multiple copies of said gene or over- expression thereof, can substantially increase the whole kernel levels of certain amino acids, such as lysine, methionine, threonine and the like. As used herein, the term "substantially increased" means that the level of a given amino acid is at least 10-20% above that present in the corresponding plant or plant part which has not been transformed with said seed storage protein gene.

In preferred embodiments, this invention produces the fertile, transgenic plants by means of a recombinant DNA-coated microprojectile bombardment of clumps of friable embryogenic callus, followed by a controlled regimen for selection of transformed callus lines.

Brief Description of the Drawings

Figure 1A shows a map of plasmid vector pHYGI1

30 utilized in Example I. Figure 1B shows the relevant part of linearized pHYGI1 encompassing the HPT coding sequence and associated regulatory elements. The base pair numbers start from the 5' nucleotide in the recognition sequence for the indicated restriction

35 enzymes, beginning with the EcoRI site at the 5' end of the CaMV 35S promoter.

Figure 2A shows a map of plasmid vector pBII221

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utilized in Example I. Figure 2B depicts linearized pBII221, e.g., from the HindIII cleavage site to the EcoRI cleavage site.

Figure 3 is a Southern blot of DNA isolated

5 from the PH1 callus line and an untransformed control
callus line, and a schematic depiction of the pHYGI1
probes used in the assay.

Figure 4 is a Southern blot of leaf DNA isolated from R0 plants regenerated from PH1 and untransformed callus, and a schematic depiction of the pHYGI1 probes used in the assay.

Figure 5 is a Southern blot of leaf DNA isolated from R1 progeny of PH1 R0 plants and untransformed R0 plants, and a schematic depiction of the pHYGI1 probes used in the assay.

Figure 6 is a Southern blot of DNA isolated from the PH2 callus line and an untransformed control callus line, and a schematic depiction of the pHYGI1 probes used in the assay.

Figure 7A shows a map of plasmid vector pZ27Z10 utilized in Example II. Figure 7B depicts linearized plasmid pZ27Z10 encompassing the z10 coding sequence and associated regulatory elements.

Figure 8 is a schematic depiction of the 25 location of the PCR primers within the chimeric z27-z10 gene.

Description of the Preferred Embodiments

The present invention is directed to the

30 production of fertile transgenic plants and seeds of the species Zea mays and to the plants, plant tissues, and seeds derived from such transgenic plants, as well as the subsequent progeny and products derived therefrom, preferably those transgenic Zea mays plants having improved food or feed value. The transgenic plants produced herein include all plants of this species, including field corn, popcorn, sweet corn, flint corn

and dent corn.

20 spontaneous mutation.

"Transgenic" is used herein to include any cell, cell line, callus, tissue, plant part or plant, the genotype of which has been altered by the presence 5 of recombinant DNA, which DNA has also been referred to in the art of genetic engineering as "heterologous DNA," "exogenous DNA" or "foreign DNA," wherein said DNA was introduced into the genotype by a process of genetic engineering, or which was initially introduced into the 10 genotype of a parent plant by such a process and is subsequently transferred to later generations by sexual crosses or asexual propagation. As used herein, "genotype" refers to the sum total of genetic material within a cell, either chromosomally, or extrachromosomally borne. Therefore, the term "transgenic" as used herein does not encompass the alteration of the genotype of Zea mays by conventional plant breeding methods or by naturally occurring events such as random cross-fertilization, viral infection or

By "heritable" is meant that the DNA is capable of transmission through at least one complete sexual cycle of a plant, i.e., it is passed from one plant through its gametes to its progeny plants.

25 The transgenic plants of this invention may be produced by (i) establishing a regenerable cell culture, preferably a friable embryogenic callus, (ii) transforming said cell culture by a microprojectile bombardment technique, (iii) identifying or selecting transformed cells, and (iv) regenerating fertile transgenic plants from the transformed cells. Some of the plants of this invention may be produced from the transgenic seed produced from the fertile transgenic plants using conventional crossbreeding techniques to develop transgenic elite lines and varieties, or commercial hybrid seed containing recombinant DNA.

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I. Plant Lines and Tissue Cultures

The cells which have been found particularly useful to produce the fertile transgenic maize plants herein are those callus cells which are regenerable, 5 both before and after undergoing a selection regimen as detailed further below. Generally, these cells will be derived from meristematic tissue which contains cells which have not yet terminally differentiated. tissue in graminaceous cereals in general and in maize, 10 in particular, comprise tissues found in juvenile leaf basal regions, immature tassels, immature embryos, and coleoptilar nodes. Preferably, immature embryos are used. Methods of preparing and maintaining callus from such tissue and plant types are well known in the art 15 and details on so doing are available in the literature, c.f. Phillips et al. (1988), the disclosure of which is hereby incorporated by reference.

The specific callus used must be able to regenerate into a fertile plant. The specific 20 regeneration capacity of particular callus is important to the success of the bombardment/selection process used herein because during and following selection, regeneration capacity may decrease significantly. It is therefore important to start with cultures that have as 25 high a degree of regeneration capacity as possible. Callus which is more than about 3 months and up to about 36 months of age has been found to have a sufficiently high level of regenerability and thus is preferred. The regenerative capacity of a particular culture may be 30 readily determined by transferring samples thereof to regeneration medium and monitoring the formation of shoots, roots, and plantlets. The relative number of plantlets arising per petri dish or per gram fresh weight of tissue may be used as a rough quantitative 35 estimate of regeneration capacity. Generally, a culture which will produce at least one plant per gram of callus tissue is preferred.

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While maize callus cultures can be initiated from a number of different plant tissues, the cultures useful herein are preferably derived from immature maize embryos which are removed from the kernels of an ear 5 when the embryos are about 1-3 mm in length. This length generally occurs about 9-14 days after pollination. Under aseptic conditions, the embryos are placed on conventional solid media with the embryo axis down (scutellum up). Callus tissue appears from the 10 scutellum after several days to a few weeks. After the callus has grown sufficiently, the cell proliferations from the scutellum may be evaluated for friable consistency and the presence of well-defined embryos. By "friable consistency" it is meant that the tissue is 15 easily dispersed without causing injury to the cells. Tissue with this morphology is then transferred to fresh media and subcultured on a routine basis about every two weeks. Sieving to reduce clumping and/or to increase cell surface area may be employed during the subculturing period.

The callus initiation media is preferably solid. In preferred embodiments, the initiation/maintenance media is typically based on the N6 salts of Chu et al. (1975) as described in Armstrong et al. (1985) or the MS salts of Murashige et al. (1962). The basal medium is supplemented with sucrose and 2,4-dichlorophenoxyacetic acid (2,4-D). Supplements such as L-proline and casein hydrolysate have been found to improve the frequency of initiation of callus cultures, morphology, and growth. The cultures are generally maintained in the dark, though low light levels may also be used. The level of synthetic hormone 2,4-D, necessary for maintenance and propagation, should be generally about 0.3 to 3.0 mg/l.

Although successful transformation and regeneration has been accomplished herein with friable embryogenic callus, this is not meant to imply that

other transformable regenerable cells, tissue, or organs cannot be employed to produce the fertile transgenic plants of this invention. The only actual requirement for the cells which are transformed is that after transformation they must be capable of regeneration of a plant containing the recombinant DNA following the particular selection or screening procedure actually used.

For example, cells grown in liquid suspension 10 culture may be used. To establish these cultures, the type II callus, after 4-6 months, is transferred to liquid growth media. Methods and references for the production of regenerable suspension cell cultures are given by C. E. Green et al. in Maize for Biological Research, Plant Molec. Biol. Assoc. (1982) at pages 367-372, R. Phillips et al., Corn and Corn Improvement, Agronomy Soc. Amer., (3d ed., 1988) at pages 345-387, and I. Vasil, Cell Culture and Somatic Cell Genetics of Plants, Vol. I, Laboratory Procedures and Their 20 Applications, Academic Press (1984) at pages 152-158. Typically, the liquid growth media for suspension cultures is of similar formulation to the solid callus induction media. ABA (abscisic acid) (10⁻⁷M) may be added to the liquid growth media to augment regenerative 25 capacity and enhance culture vitality. It is preferred that the callus not be sieved prior to introduction into the liquid media.

The cultures in liquid media are subcultured as appropriate for maintaining active growth and their regenerative properties. In preferred embodiments, the cultures are subcultured once a week at a 1:8-9 dilution with fresh growth medium.

II. DNA Used for Transformation

As used herein, the term "recombinant DNA" refers to DNA that has been derived or isolated from any source, that may be subsequently chemically altered, and

later introduced into Zea mays. An example of recombinant DNA "derived" from a source, would be a DNA sequence that is identified as a useful fragment within a given organism, and which is then chemically synthesized in essentially pure form. An example of such DNA "isolated" from a source would be a useful DNA sequence that is excised or removed from said source by chemical means, e.g., by the use of restriction endonucleases, so that it can be further manipulated, 0 e.g., amplified, for use in the invention, by the methodology of genetic engineering.

Therefore "recombinant DNA" includes completely synthetic DNA, semi-synthetic DNA, DNA isolated from biological sources, and DNA derived from introduced RNA.

15 Generally, the recombinant DNA is not originally resident in the Zea mays genotype which is the recipient of the DNA, but it is within the scope of the invention to isolate a gene from a given Zea mays genotype, and to subsequently introduce multiple copies of the gene into the same genotype, e.g., to enhance production of a given gene product such as a storage protein.

The recombinant DNA includes but is not limited to, DNA from plant genes, and non-plant genes such as those from bacteria, yeasts, animals or viruses;

25 modified genes, portions of genes, chimeric genes, including genes from the same or different Zea mays genotype.

The recombinant DNA used for transformation herein may be circular or linear, double-stranded or 30 single-stranded. Generally, the DNA is in the form of chimeric DNA, such as plasmid DNA, that can also contain coding regions flanked by regulatory sequences which promote the expression of the recombinant DNA present in the resultant corn plant. For example, the recombinant 35 DNA may itself comprise or consist of a promoter that is active in Zea mays, or may utilize a promoter already present in the Zea mays genotype that is the trans-

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formation target.

The compositions of, and methods for, constructing recombinant DNA which can transform certain plants are well known to those skilled in the art, and 5 the same compositions and methods of construction may be utilized to produce the DNA useful herein. The specific composition of the DNA is not central to the present invention and the invention is not dependent upon the composition of the specific recombinant DNA used. 10 K. Weising et al., <u>Ann. Rev. Genetics</u>, <u>22</u>, 421 (1988) describes suitable DNA components, selectable marker genes, reporter genes, enhancers, introns, and the like, as well as provides suitable references for compositions therefrom. J. Sambrook et al., Molecular Cloning: A 15 Laboratory Manual, Cold Spring Harbor Laboratory Press (2d ed., 1989), provides suitable methods of construction. Generally, the recombinant DNA will be relatively small, i.e., less than about 30 kb to minimize any susceptibility to physical, chemical, or 20 enzymatic degradation which is known to increase as the size of the DNA increases.

Suitable recombinant DNA for use herein includes all DNA which provides for, or enhances, a beneficial feature of the resultant transgenic corn plant. The DNA may encode proteins or antisense RNA transcripts in order to promote increased food values, higher yields, pest resistance, disease resistance, herbicide resistance, and the like. For example, the DNA can encode a DHDP synthase, as does the dap A gene, for increased lysine production; Bacillus thuringiensis (Bt), δ-endotoxin or a protease inhibitor for insect resistance; bacterial EPSP synthase for resistance to glyphosate herbicide; and chitinase or glucan endo-1,3-β-glucosidase for fungicidal properties.

Of importance in improving food or feed value are genes encoding proteins that contain high levels of essential amino acids. For example, to be nutritionally

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adequate and support optimal growth of chickens, corn-soybean meal poultry feed is generally supplemented with synthetic methionine or a methionine analog. The development of lines of corn which supply higher levels of methionine can reduce the need for methionine supplements. The development of such high methionine corn lines can be accomplished by introducing into the corn genome a highly expressed gene or genes encoding a high-methionine protein.

Examples of genes encoding high-methionine 10 proteins include: 1.) the gene encoding a maize 15 kDzein protein (11% methionine), (Pedersen et al., J. Biol. Chem., 261, 6279 (1986)); 2.) the gene encoding a Brazil nut storage protein (18% methionine), (Altenbach 15 et al., Plant Mol. Biol., 8; 239 (1987)) and 3.) the gene encoding a maize 10 kD-zein protein (22.5% methionine), (Kirihara et al., Gene, 7, 359 (1988)). The preferred gene is the 10 kD-zein gene, since it is an endogenous maize gene whose protein product normally · 20 accumulates in the kernel and is twice as high in methionine than the 15 kD-zein protein. To obtain high levels of expression in the seed, its coding sequence optionally may be fused to the regulatory sequence from a highly-expressed, seed-specific gene. Alternatively, introduction of additional copies of the intact endogenous 10 kD-zein gene into the corn genome can also increase the methionine content of corn seed.

Lysine, an amino acid essential in the diets of humans and monogastric animals, is among the three most limiting amino acids in most of the staple crops, the cereals in particular. Consequently, grain-based diets must be supplemented with synthetic lysine or with lysine-containing oilseed protein meals. Further, since most oilseed meals are themselves inadequate lysine sources, balancing the feed mixture for lysine frequently results in meals which are too high in other, less desirable nutrients. Therefore, a method to

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increase the lysine content of either the cereal grains or the oilseed crops or both would result in significant added nutritional value, as well as a significant cost savings to end-users such as the swine and poultry producers.

One approach to improving the lysine content of cereals is to deregulate the biosynthetic pathway, allowing free lysine to accumulate. The dap A gene of Escherichia coli encodes dihydrodipicolinic acid

10 synthase (DHDPS), a key regulatory enzyme whose activity in plants is strongly feedback-inhibited by lysine. The bacterial enzyme is about 200-fold less sensitive to inhibition by lysine. The introduction and expression of the dap A gene in plant cells would allow the

15 synthesis of free lysine to continue after the native plant DHDPS has been completely inhibited.

Of particular importance in maintaining yield from corn plants and contributing significantly to controlling the cost of growing a corn crop is the 20 protection of the corn against attack by insect pests. In the USA the major insect pests of corn include a variety of Lepidoptera pests such as the European corn borer, cutworms and earworms as well as Coleoptera species such as Diabrotica spp. The protection of corn 25 against insect attack is expensive to the grower and requires the use of toxic chemical insecticides applied in a timely manner. Since traditional methods of breeding and selection have not allowed for the development of new lines of corn that are substantially 30 resistant to major insect pests, the introduction and inheritance of insect resistance genes or sequences in corn plants in accord with the present invention would reduce costs to the grower, reduce the use of toxic chemical insecticides and provide for more effective 35 control of insect pests.

Essential components of the present invention are the introduction of the insect resistance gene into

the corn cell, mitotic replication of the gene so that the gene is incorporated into whole corn plants and is ultimately inherited by subsequent offspring of the plant via the processes of mitotic and meiotic division.

include nearly 20 known subspecies of bacteria which produce endotoxin polypeptides that are toxic when ingested by a wide variety of insect species. The biology and molecular biology of the endotoxin proteins (Bt proteins) and corresponding genes (Bt genes) has been reviewed recently by H.R. Whitely et al., Ann. Rev. Microbiol., 40, 549 (1986) and by H. Hofte et al., Microbiol. Rev., 53, 242 (1989). Genes coding for a variety of Bt proteins have been cloned and sequenced.

Research has demonstrated that a segment of the Bt polypeptide is essential for toxicity to a variety of Lepidoptera pests and is contained within approximately the first 50% of the Bt polypeptide molecule.

Consequently, a truncated Bt polypeptide coded by a

truncated Bt gene will in many cases retain its toxicity towards a number of Lepidoptera insect pests. The HD73 and HD1 Bt polypeptides have been shown to be toxic to the larvae of the important Lepidoptera insect pests of corn plants in the USA such as the European corn borer,

25 cutworms and earworms. The genes coding for the HD1 and HD73 Bt polypeptides have been cloned and sequenced by M. Geiser et al., Gene, 48, 109 (1986) and M.J. Adang et al., Gene, 36, 289 (1985), respectively, and can be cloned from HD1 and HD73 strains obtained from culture collections (e.g. Bacillus Genetic Stock Center,

Collections (e.g. Bacillus Genetic Stock Center, Columbus, Ohio or USDA Bt stock collection Peoria, Illinois) using standard protocols.

DNA coding for new, previously uncharacterized Bt toxins, may be cloned from the host Bacillus organism using protocols that have previously been used to clone Bt genes. These include the construction of a bank of DNA isolated from the Bacillus organism in a suitable

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plasmid or phage vector replicated in a suitable host and the use of antibodies, raised against the Bt protein or DNA isolated from a homologous Bt gene sequence, to identify transformants that contain the cloned Bt 5 sequence. The approximate location of the Bt coding sequence may be initially determined using deletion analysis of the cloned DNA. The precise location of the Bt coding sequence could be determined using a variety of standard methods, including the determination of the 10 sequence of the cloned segment of DNA, determining the presence of large open reading frames in this sequence that could code for the Bt protein and confirmation of the Bt coding nature of the DNA sequence by comparing the amino acid sequence derived from the DNA sequence 15 with that derived from partial amino acid sequencing of the Bt protein.

A chimeric Bt gene useful in the present invention would comprise a 5' DNA sequence, comprising a sequence of DNA which will allow for the initiation of 20 transcription ("promoter") and translation of a downstream located Bt sequence in a corn plant. chimeric Bt gene would also comprise a 3' DNA sequence that includes a sequence derived from the 3' non-coding region of a gene that can be expressed in corn. 25 importantly, the chimeric Bt gene will include a DNA sequence coding for a toxic Bt polypeptide produced by Bacillus thuringiensis or toxic portions thereof or having substantial amino sequence homology thereto. Bt coding sequence would include: (i) DNA sequences 30 which code for insecticidally active proteins that have substantial homology to Bt endotoxins that are active against insect pests of corn, e.g., the HD73 or HD1 Bt sequences; (ii) sequences coding for insecticidally active segments of the Bt endotoxin polypeptide, e.g., 35 insecticidally active HD73 or HD1 polypeptides truncated from the carboxy and/or amino termini; (iii) a truncated Bt sequence fused in frame with a sequence(s) that codes

for a polypeptide that provides some additional advantage such as: (a) genes that are selectable, e.g., genes that confer resistance to antibiotics or herbicides, (b) reporter genes whose products are easy to detect or assay, e.g., luciferase or betaglucuronidase; (c) DNA sequences that code for polypeptide sequences that have some additional use in stabilizing the Bt protein against degradation or enhance the efficacy of the Bt protein against insects, e.g., protease inhibitors and (d) sequences that help direct the Bt protein to a specific compartment inside or outside the corn cell, e.g., a signal sequence.

To obtain optimum synthesis of the Bt protein in corn, it may also be appropriate to adjust the DNA 15 sequence of the Bt gene to more resemble the genes that are efficiently expressed in corn. Since the codon usage of many Bt genes, including the HD73 and HD1 genes, is more similar to that used by Bacillus species and dissimilar to that used by genes that are expressed 20 in maize, the expression of the Bt gene in maize cells can be improved by the replacement of these rarely used Bacillus codons with those that are used more frequently in maize plants (See E. Murray et al., Nucl. Acids Res., 17, 477 (1989)). Such replacement of codons would 25 require the substitution of bases without changing the amino acid sequence of the resulting Bt polypeptide. The Bt polypeptide would be identical in sequence to the bacterial gene or segments thereof. The complete Bt coding sequence, or sections thereof, containing a 30 higher proportion of maize preferred codons than the original bacterial gene could be synthesized using standard chemical synthesis protocols, and introduced or assembled into the Bt gene using standard protocols, such as site-directed mutagenesis or DNA polymerization 35 and ligation and the like.

Aside from recombinant DNA sequences that serve as transcription units or portions thereof, useful

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recombinant DNA may be untranscribed, serving a regulatory or a structural function. Also, the DNA may be introduced to act as a genetic tool to generate mutants and/or assist in the identification, genetic tagging, or isolation of segments of corn DNA. Additional examples may be found in Weising, cited supra.

The recombinant DNA to be introduced into the plant cells further will generally contain either a selectable marker or a reporter gene or both to

10 facilitate identification and selection of transformed cells. Alternatively, the selectable marker may be carried on a separate piece of DNA and used in a co-transformation procedure. Both selectable markers and reporter genes may be flanked with appropriate

15 regulatory sequences to enable expression in plants. Useful selectable markers are well known in the art and include, for example, antibiotic and herbicide resistance genes.

Specific examples of selectable marker genes 20 are disclosed in Weising et al., cited supra. A preferred selectable marker gene is the hygromycin phosphotransferase (HPT) coding sequence, which may be derived from E. coli and which confers resistance to the antibiotic hygromycin B. Other selectable markers 25 include the aminoglycoside phosphotransferase gene of transposon Tn5 (AphII) which encodes resistance to the antibiotics kanamycin, neomycin and G418, as well as those genes which code for resistance or tolerance to glyphosate, 2,2-dichloropropionic acid, methotrexate, 30 imidazolinone herbicides, sulfonylurea herbicides, bromoxynil, phosphinothricin and other herbicidal compounds. Those selectable marker genes which confer resistance or tolerance to these phytotoxic compounds are also of commercial utility in the resulting 35 transformed plants. Selectable marker genes encoding

enzymes which impart resistance to phytotoxic compounds are listed in Table 1, below.

Table 1

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Selectable Marker Genes

10	Resistance Gene or Enzyme	Confers Resistance to:	Reference
	Neomycin phospho- transferase (neo)	G-418, neomycin, kanamycin	P.J. Southern et al., <u>J.</u> <u>Mol. Appl. Gen.</u> , <u>1</u> , 327 (1982)
15	Hygromycin phos- photranferase (hpt or hyg)	Hygromycin B	Y. Shimizu et al., Mol. Cell Biol., 6, 1074 (1986)
20	Dihydrofolate reductase (dhfr)	Methotrexate	W.W. Kwok et al., <u>PNAS USA</u> , 4552 (1986)
25	Phosphinothricin acetyltransferase (bar)	Phosphinothricin	M. DeBlock et al., EMBO J., <u>6</u> , 2513 (1987)
25	2,2-Dichloropro- pionic acid dehalogenase	2,2-Dichloropro- pionic acid (Dalapon)	V. Buchanan-Wollaston et al., J. Cell. Biochem., Supp. 13D, 330 (1989)
30	Acetohydroxyacid synthase	Sulfonylurea, imidazolinone and triazolopyrimidine herbicides	P.C. Anderson et al. (U.S. Patent No. 4,761,373); G.W. Haughn et al., Mol. Gen. Genet., 211, 266 (1988)
35	5-Enolpyruvyl- shikimate-phosphate synthase (aroA)	Glyphosate	L. Comai et al., <u>Nature</u> , <u>317</u> , 741 (1985)
40	Haloarylnitrilase	Bromoxynil	D.M. Stalker et al., published PCT appln. W087/04181
45	Acetyl-coenzyme A carboxylase	Sethoxydim, haloxyfop	W.B. Parker et al., <u>Plant</u> <u>Physiol.</u> , <u>92</u> , 1220 (1990)
	Dihydropteroate synthase (sul I)	Sulfonamide herbicides	F. Guerineau et al., <u>Plant Molec. Biol.</u> , <u>15</u> , 127 (1990)
50	32 kD photosystem II polypeptide (psbA)	Triazine herbicides	J. Hirschberg et al., <u>Science</u> , <u>222</u> , 1346 (1983)

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Anthranilate synthase

5-Methyltryptophan

K. Hibberd et al., (U.S. Patent No. 4,581,847)

Dihydrodipicolinic 5 acid synthase (dap A)

Dihydrodipicolinic Aminoethyl cysteine

K. Glassman et al., published PCT application No. W089/11789

Reporter genes are used for identifying potentially transformed cells and for evaluating the 5 functionality of regulatory sequences. Reporter genes which encode for easily assayable marker proteins are well known in the art. In general, a reporter gene is a gene which is not present in or expressed by the recipient organism or tissue and which encodes a protein 10 whose expression is manifested by some easily detectable property, e.g., phenotypic change or enzymatic activity. Examples of such genes are provided in Weising et al., supra. Preferred genes include the chloramphenicol acetyl transferase gene (cat) from Tn9 of E. coli, the 15 beta-glucuronidase gene (gus) of the \underline{uidA} locus of \underline{E} . coli, and the luciferase gene from firefly Photinus pyralis. Expression of the reporter gene is assayed at a suitable time after the DNA has been introduced into the recipient cells. A preferred such assay entails the 20 use of the E. coli beta-glucuronidase (GUS) gene (R. Jefferson et al., <u>EMBO J.</u>, <u>16</u>, 3901 (1987)). Maize cells transformed and expressing this gene will stain blue upon exposure to the substrate, 5-bromo-4-chloro-3indolyl-beta-D-glucuronide (X-GLUC), in the 25 extracellular medium.

The regulatory sequences useful herein include any constitutive, inducible, tissue or organ specific, or developmental stage specific promoter which can be expressed in the particular plant cell. Suitable such promoters are disclosed in Weising et al., supra. The following is a partial representative list of promoters

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suitable for use herein: regulatory sequences from the T-DNA of Agrobacterium tumefaciens, including mannopine synthase, nopaline synthase, and octopine synthase; alcohol dehydrogenase promoter from corn; light-inducible promoters such as the ribulose-bisphosphate-carboxylase/oxygenase small subunit genes from a variety of species; and the major chlorophyll a/b binding protein gene promoters; 35S and 19S promoters of cauliflower mosaic virus (CaMV); developmentally regulated promoters such as the waxy, zein, or bronze promoters from maize; as well as synthetic or other natural promoters which are either inducible or constitutive, including those promoters exhibiting organ-specific expression or expression at specific development stage(s) of the plant.

Other elements such as introns, enhancers, polyadenylation sequences and the like, may also be a part of the recombinant DNA. Such elements may or may not be necessary for the function of the DNA, but may 20 provide improved expression of the DNA by affecting transcription, stability of the mRNA, or the like. Such elements may be included in the DNA as desired to obtain the optimal performance of the transforming DNA in the plant. For example, the maize AdhIS first intron may be 25 placed between the promoter and the coding sequence in a particular recombinant DNA construction. This intron, when included in a DNA construction, is known to increase production of a protein in maize cells. (J. Callis et al., Genes and Develop., 1, 1183 (1987)). 30 However, sufficient expression for a selectable marker to perform satisfactorily can often be obtained without an intron. (T. Klein et al., Plant Physiol., 91, 440 (1989)). An example of an alternative suitable intron is the shrunken-1 first intron of Zea mays. These other 35 elements must be compatible with the remainder of the DNA constructions.

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III. DNA Delivery Process

The recombinant DNA can be introduced into the regenerable maize cell cultures, preferably into callus cultures via a particle bombardment process. A general description of a suitable particle bombardment instrument is provided in Sanford et al. (1987), the disclosure of which is incorporated herein by reference. While protocols for the use of the instrument in the bombardment of maize non-regenerable suspension culture cells are described in Klein et al. (1988a, 1988b, and 1989), no protocols have been published for the bombardment of callus cultures or regenerable maize cells.

In a microprojectile bombardment process, also referred to as a biolistic process, the transport of the recombinant DNA into the callus is mediated by very small particles of a biologically inert material. When the inert particles are coated with DNA and accelerated to a suitable velocity, one or more of the particles is able to enter into one or more of the cells where the DNA is released from the particle and expressed within the cell. Some of the recipient cells stably retain the introduced DNA and express it.

The particles, called microprojectiles, are

generally of a high density material such as tungsten or
gold. They are coated with the DNA of interest. The
microprojectiles are then placed onto the surface of a
macroprojectile which serves to transfer the motive
force from a suitable energy source to the

microprojectiles. After the macroprojectile and the
microprojectiles are accelerated to the proper velocity,
they contact a blocking device which prevents the
macroprojectile from continuing its forward path but
allows the DNA-coated microprojectiles to continue on
and impact the recipient callus cells. Suitable such
instruments may use a variety of motive forces such as
quipowder or shock waves from an electric arc discharge

(P. Christou et al., <u>Plant Physiol.</u>, <u>87</u>, 671 (1988)). An instrument in which gunpowder is the motive force is currently preferred and such is described and further explained in Sanford et al. (1987), the disclosure of which is incorporated herein by reference.

A protocol for the use of the gunpowder instrument is provided in Klein et al. (1988a,b) and involves two major steps. First, tungsten microprojectiles are mixed with the DNA, calcium 10 chloride, and spermidine in a specified order in an aqueous solution. The concentrations of the various components may be varied as taught. The preferred procedure entails exactly the procedure of Klein et al. (1988b) except for doubling the stated optimum DNA 15 concentration. Secondly, the DNA-coated microprojectiles, macroprojectiles, and recipient cells are placed in position in the instrument and the motive force is applied to the macroprojectiles. Parts of this step which may be varied include the distance of the 20 recipient cells from the end of the barrel as well as the vacuum level in the sample chamber. The recipient tissue is positioned 2-15, preferably 5 cm below the stopping plate tray.

25 generation of transgenic plants should generally be about midway between transfer periods, and thus, past any "lag" phase that might be associated with a transfer to a new media, but also before reaching any "stationary" phase associated with an extended period of time without subculture to fresh media. The tissue may be used in the form of pieces of about 30 to 80, preferably about 40 to 60, mg. The clumps can be placed on a petri dish or other surface and arranged in essentially any manner, recognizing that (i) the space in the center of the dish will receive the heaviest concentration of metal-DNA particles and the tissue located there is likely to suffer damage during bombardment and,

(ii) the number of particles reaching the tissue will decrease with increasing distance of the tissue from the center of the blast so that the tissue far from the center of the dish is less likely to be bombarded. A 5 mesh screen, preferably of metal, may be laid on the dish to prevent splashing or ejection of the tissue. An alternative method for presentation of the tissue for bombardment is to spread the tissue onto a filter paper in a thin layer. The tissue may be bombarded one or 10 more times with the DNA-coated metal particles.

IV. Selection Process

Once the calli have been bombarded with the recombinant DNA and the DNA has penetrated some of the 15 cells, it is necessary to identify and select those cells which both contain the recombinant DNA and still retain sufficient regenerative capacity. There are two general approaches which have been found useful for accomplishing this. First, the transformed calli or 20 plants regenerated therefrom can be screened for the presence of the recombinant DNA by various standard methods which could include assays for the expression of reporter genes or assessment of phenotypic effects of the recombinant DNA, if any. Alternatively, and 25 preferably, when a selectable marker gene has been transmitted along with or as part of the recombinant DNA, those cells of the callus which have been transformed can be identified by the use of a selective agent to detect expression of the selectable marker 30 gene.

Selection of the putative transformants is a critical part of the successful transformation process since selection conditions must be chosen so as to allow growth and accumulation of the transformed cells while simultaneously inhibiting the growth of the non-transformed cells. The situation is complicated by the fact that the vitality of individual cells in a

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population is often highly dependent on the vitality of neighboring cells. Also, the selection conditions must not be so severe that the plant regeneration capacity of the callus cells and the fertility of the resulting

5 plant are precluded. Thus, the effects of the selection agent on cell viability and morphology should be evaluated. This may be accomplished by experimentally producing a growth inhibition curve for the given selective agent and tissue being transformed beforehand.

10 This will establish the concentration range which will inhibit growth.

When a selectable marker gene has been used, the callus clumps may be either allowed to recover from the bombardment on non-selective media, or preferably, directly transferred to media containing the selection agent.

Selection procedures involve exposure to a toxic agent and may employ sequential changes in the concentration of the agent and multiple rounds of 20 selection. The particular concentrations and cycle lengths are likely to need to be varied for each particular agent. A currently preferred selection procedure entails using an initial selection round at a relatively low toxic agent concentration and then later round(s) at higher concentration(s). This allows the selective agent to exert its toxic effect slowly over a longer period of time. Preferably, the concentration of the agent is initially such that about a 5-40% level of growth inhibition will occur, as determined from a growth inhibition curve. The effect may be to allow the 30 transformed cells to preferentially grow and divide while inhibiting untransformed cells, but not to the extent that growth of the untransformed cells is prevented. Once the few individual transformed cells 35 have divided a sufficient number of times, the tissue may be shifted to media containing a higher concentration of the toxic agent to kill essentially all

untransformed cells. The shift to the higher concentration also reduces the possibility of non-transformed cells adapting to the agent. The higher level is preferably in the range of about 30 to 100% growth inhibition. The length of the first selection cycle may be from about 1 to 4 weeks, preferably about 2 weeks. Later selection cycles may be from about 1 to 12 weeks, preferably about 2 to 10 weeks. Putative maize transformants can generally be identified as proliferating sectors of tissue among a background of non-proliferating cells. The callus may also be cultured on non-selective media at various times during the overall selection procedure.

Once a callus sector is identified as a 15 putative transformant, transformation can be confirmed by phenotypic and/or genotypic analysis. If a selection agent is used, an example of phenotypic analysis is to measure the increase in fresh weight of the putative transformant as compared to a control on various levels 20 of the selective agent. Other analyses that may be employed will depend on the function of the recombinant DNA. For example, if an enzyme or protein is encoded by the DNA, enzymatic or immunological assays specific for the particular enzyme or protein may be used. Other 25 gene products may be assayed by using a suitable bioassay or chemical assay. Other such techniques are well known in the art and are not repeated here. presence of the gene can also be confirmed by conventional procedures, i.e., Southern blot or 30 polymerase chain reaction (PCR) or the like.

V. Regeneration of Plants and Production of Seed

Cell lines which have been shown to be
transformed must then be regenerated into plants and the

fertility of the resultant plants determined.

Transformed lines which test positive by genotypic
and/or phenotypic analysis are then placed on a medium

which promotes tissue differentiation and plant regeneration. Regeneration may be carried out in accordance with standard procedures well known in the art. The procedures commonly entail reducing the level of auxin which discontinues proliferation of a callus and promotes somatic embryo development or other tissue differentiation. One example of such a regeneration procedure is described in Green et al. (1982). The plants are grown to maturity in a growth room or greenhouse and appropriate sexual crosses are made as described by Neuffer (1982).

Regeneration, while important to the present invention, may be performed in any conventional manner. If a selectable marker has been transformed into the cells, the selection agent may be incorporated into the regeneration media to further confirm that the regenerated plantlets are transformed. Since regeneration techniques are well known and not critical to the present invention, any technique which accomplishes the regeneration and produces fertile plants may be used.

VI. Analysis of R1 Progeny

The plants regenerated from the transformed

25 callus are referred to as the R0 generation or R0

plants. The seeds produced by various sexual crosses of
the R0 generation plants are referred to as R1 progeny
or the R1 generation. When R1 seeds are germinated, the
resulting plants are also referred to as the R1

30 generation.

To confirm the successful transmission and inheritance of the recombinant DNA through one complete sexual cycle, the Rl generation should be analyzed to confirm the presence of the transforming DNA. The analysis may be performed in any of the manners such as were disclosed above for analyzing the bombarded callus for evidence of transformation, taking into account the

fact that plants and plant parts are being used in place of the callus.

The recombinant DNA can show different types of inheritance patterns for different transformed lines.

5 For example, the recombinant DNA may be inherited in the progeny according to the rules of Mendelian inheritance. This type of heritability involves transmission of the DNA through both the male and female gametes and is associated with stable integration or incorporation into 10 maize nuclear DNA. An alternative type of inheritance pattern is maternal inheritance, in which the recombinant DNA is transmitted primarily or exclusively through the female gametes. The inheritance pattern for a particular transformant may be ascertained by analysis of the progeny of various sexual crosses.

VII. Establishment of the Recombinant DNA in Other Maize Varieties

Fertile, transgenic plants may then be used in 20 a conventional maize breeding program in order to incorporate the introduced recombinant DNA into the desired lines or varieties. Methods and references for convergent improvement of corn are given by Hallauer et al., (1988) incorporated herein by reference. Among the 25 approaches that conventional breeding programs employ is a conversion process (backcrossing). Briefly, conversion is performed by crossing the initial transgenic fertile plant to elite inbred lines. progeny from this cross will segregate such that some of 30 the plants will carry the recombinant DNA whereas some will not. The plants that do carry the DNA are then crossed again to the elite inbred lines resulting in progeny which segregate once more. This backcrossing process is repeated until the original elite inbred has 35 been converted to a line containing the recombinant DNA, yet possessing all important attributes originally found in the parent. Generally, this will require about 6-8

generations. A separate backcrossing program will be generally used for every elite line that is to be converted to a genetically engineered elite line.

Generally, the commercial value of the

transformed corn produced herein will be greatest if the
recombinant DNA can be incorporated into many different
hybrid combinations. A farmer typically grows several
hybrids based on differences in maturity, standability,
and other agronomic traits. Also, the farmer must

select a hybrid based upon his or her geographic
location since hybrids adapted to one region are
generally not adapted to another because of differences
in such traits as maturity, disease, and insect
resistance. As such, it is necessary to incorporate the
recombinant DNA into a large number of parental lines so
that many hybrid combinations can be produced containing
the desirable heterologous DNA.

Corn breeding and the techniques and skills required to transfer genes from one line or variety to another are well known to those skilled in the art.

Thus, introducing recombinant DNA into any other line or variety can be accomplished by these breeding procedures.

25 <u>VIII. Uses of Transgenic Plants</u>

The transgenic plants produced herein are expected to be useful for a variety of commercial and research purposes. Transgenic plants can be created for use in traditional agriculture to possess traits

30 beneficial to the grower (e.g., agronomic traits such as pest resistance, herbicide resistance or increased yield), beneficial to the consumer of the grain harvested from the plant (e.g., improved nutritive content in human food or animal feed), or beneficial to the food processor (e.g., improved processing traits). In such uses, the plants are generally grown for the use of their grain in human or animal foods. However, other

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parts of the plants, including stalks, husks, vegetative parts, and the like, may also have utility, including use as part of animal silage or for ornamental purposes. Often, chemical constituents (e.g., oils or starches) of 5 corn and other crops are extracted for foods or industrial use and transgenic plants may be created which have enhanced or modified levels of such components.

Transgenic plants may also find use in the 10 commercial manufacture of proteins or other molecules, where the molecule of interest is extracted or purified from plant parts, seeds, and the like. Cells or tissue from the plants may also be cultured, grown in vitro, or fermented to manufacture such molecules.

The transgenic plants may also be used in commercial breeding programs, or may be crossed or bred to plants of related crop species. Improvements encoded by the recombinant DNA may be transferred, e.g., from corn cells to cells of other species, e.g., by 20 protoplast fusion.

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The transgenic plants may have many uses in research or breeding, including creation of new mutant plants through insertional mutagenesis, in order to identify beneficial mutants that might later be created 25 by traditional mutation and selection. An example would be the introduction of a recombinant DNA sequence encoding a transposable element that may be used for generating genetic variation. The methods of the invention may also be used to create plants having 30 unique "signature sequences" or other marker sequences which can be used to identify proprietary lines or varieties.

The following non-limiting examples are illustrative of the present invention. 35 presented to better explain the general procedures which were used to prepare the fertile Zea mays plants of this invention which stably express the recombinant DNA and

which transmit that DNA to progeny. All parts and percentages are by weight unless otherwise specified. It must be recognized that the probability of a specific transformation event occurring is a function of the amount of material subjected to the transformation procedure. Thus, when individual situations arise in which the procedures described herein do not produce a transformed product, repetition of the procedures will be required.

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EXAMPLE I

Fertile Transgenic Zea Mays Plants from Callus Line AB11

15 <u>I. Initiation and maintenance of maize cell cultures</u>
which retain plant regeneration capacity

Friable, embryogenic maize callus cultures were initiated from hybrid immature embryos produced by pollination of inbred line A188 plants (University of .20 Minnesota, Crop Improvement Association) with pollen of inbred line B73 plants (Iowa State University). Ears were harvested when the embryos had reached a length of 1.5 to 2.0 mm. Each ear was surface sterilized in 50% v/v commercial bleach (2.63% w/v sodium hypochlorite) 25 for 20 min. at room temperature. The ears were then washed with sterile, distilled, deionized water. Immature embryos were aseptically isolated and placed on nutrient initiation/maintenance medium with the root/shoot axis exposed to the medium. Initiation/ 30 maintenance medium (hereinafter referred to as "F medium") consisted of N6 basal media (Chu 1975) with 2% (w/v) sucrose, 1.5 mg per liter 2,4-dichlorophenoxyacetic acid (2,4-D), 6 mM proline, and 0.25% Gelrite (Kelco, Inc., San Diego). The pH was adjusted to 5.8 35 prior to autoclaving. Unless otherwise stated, all tissue culture manipulations were carried out under sterile conditions.

The immature embryos were incubated at 26°C in

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the dark. Cell proliferations from the scutellum of the immature embryos were evaluated for friable consistency and the presence of well-defined somatic embryos. Tissue with this morphology was transferred to fresh 5 media 10 to 14 days after the initial plating of the immature embryos. The tissue was then subcultured on a routine basis every 14 to 21 days. Sixty to eighty milligram quantities of tissue were removed from pieces of tissue that had reached a size of approximately one 10 gram and were transferred to fresh media. Subculturing always involved careful visual monitoring to be sure that only tissue of the correct morphology was maintained. The presence of somatic embryos ensured that the cultures would give rise to plants under the 15 proper conditions. The cell culture named AB11 used in this example was such a culture and had been initiated about 1 year before bombardment.

II. Plasmids

The plasmid pHYGI1, was constructed in the vector pBS+ (Stratagene, Inc., San Diego, CA), a 3.2 Kb circular plasmid, using standard recombinant DNA techniques. The 553 bp Bcl-BamHI fragment containing the maize AdhIS first intron (Callis et al. 1987) was inserted between the CaMV 35S promoter and the hygromycin coding sequence of pCHN1-1, a plasmid constructed in accord with Example V. A map of pHYGI1 is provided as Figure 1. A sample of pHYGI1 was deposited at the American Type Culture Collection,

Rockville, MD, USA, on March 16, 1990, under the provisions of the Budapest Treaty, and assigned accession number 40774.

pBII221 contains the <u>E</u>. <u>coli</u> β-glucuronidase coding sequence flanked at the 5' end by the CaMV 35S promoter and at the 3' end by the nos polyadenylation sequence. The plasmid was constructed by inserting the maize AdhIS first intron between the 35S promoter and

the coding sequence of pBI221 (Jefferson et al. 1987). A map of pBII221 is provided as Figure 2.

Plasmids were introduced into the embryogenic callus culture AB11 by microprojectile bombardment.

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III. DNA delivery process

The embryogenic maize callus line AB11 was subcultured 7 to 12 days prior to microprojectile bombardment. AB11 callus was prepared for bombardment as follows. Five clumps of callus, each approximately 50 mg in wet weight were arranged in a cross pattern in the center of a sterile 60 x 15 mm petri plate (Falcon 1007). Plates were stored in a closed container with moist paper towels, throughout the bombardment process. Twelve plates were prepared.

Plasmids were coated onto M-10 tungsten particles (Biolistics) exactly as described by Klein et al. (1988b) except that, (i) twice the recommended quantity of DNA was used, (ii) the DNA precipitation onto the particles was performed at 0°C, and (iii) the tubes containing the DNA-coated tungsten particles were stored on ice throughout the bombardment process.

All of the tubes contained 25 µl of 50 mg/ml M10 tungsten in water, 25 µl of 2.5 M CaCl₂, and 10 µl of
25 100 mM spermidine along with a total of 5 µl of 1 mg/ml
plasmid DNA. When both plasmids were used, each was
present in an amount of 2.5 µl. One tube contained only
plasmid pBII221; and one tube contained T.E. buffer (see
Table 2, below).

All tubes were incubated on ice for 10 min., the particles were pelleted by centrifugation in an Eppendorf centrifuge at room temperature for 5 seconds, 25 µl of the supernatant was discarded. The tubes were stored on ice throughout the bombardment process. Each preparation was used for no more than 5 bombardments.

Macroprojectiles and stopping plates were obtained from Biolistics, Inc. (Ithaca, NY). They were

sterilized as described by the supplier. The microprojectile bombardment instrument was obtained from Biolistics, Inc.

The sample plate tray was placed 5 cm below the bottom of the stopping plate tray of the microprojectile instrument, with the stopping plate in the slot nearest to the barrel. Plates of callus tissue prepared as described above were centered on the sample plate tray and the petri dish lid removed. A 7 x 7 cm square rigid wire mesh with 3 x 3 mm mesh and made of galvanized steel was placed over the open dish in order to retain the tissue during the bombardment. Tungsten/DNA preparations were sonicated as described by Biolistics, Inc. and 2.5 µl of the suspensions were pipetted onto the top of the macroprojectiles for each bombardment. The instrument was operated as described by the manufacturer. The bombardments which were performed are summarized on Table 2.

20		Table 2						
		2 x pBII221 prep To determine transient expression frequency						
25		7 x pHYGI1/pBII221 (1:1) As a potential positive treatment for transformation						
30		3 x T.E.ª Negative control treatment						
	а	10mM Tris·HCL, pH 8.0, 1 mM EDTA						

³⁵ The two plates of callus bombarded with pBII221 were transferred plate for plate to F medium (with no hygromycin) and the callus cultured at 26°C in the dark. After 2 days, this callus was then transferred plate for plate into 35 x 10 mm petri plates (Falcon 1008) containing 2 ml of GUS assay buffer (1 mg/ml of 5-bromo-4-chloro-3-indolyl-beta-D-glucuronide) (Research Organics), 100 mM sodium phosphate pH 7.0, 5 mM each of

potassium ferricyanide and potassium ferrocyanide, 10 mM EDTA, and 0.06% Triton X-100. The plates were incubated at 37°C for 3 days after which the number of blue cells was counted giving a total of 313 and 355 transient GUS-expressing cells visible in the two plates, suggesting that the DNA delivery process had also occurred with the other bombarded plates. The plates of tissue used in the GUS assay were discarded after counting since the GUS assay is destructive.

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IV. Selection process

Hygromycin B (Calbiochem) was incorporated into the medium prior to pouring plates by addition of the appropriate volume of filter-sterilized 100 $\rm mg/ml$

15 hygromycin B dissolved in water, when the medium had cooled to 45°C.

Immediately after all samples had been bombarded, callus from all of the plates treated with pHYGI1/pBII221, and two of the T.E. plates was

20 transferred plate for plate onto F medium containing 15 mg/l hygromycin B, (ten pieces of callus per plate). These are referred to as round 1 selection plates.

Callus from the T.E. treated plate was transferred to F

medium without hygromycin. This tissue was subcultured 25 every 2-3 weeks onto nonselective medium and is referred to as unselected control callus.

After 14 days of selection, tissue appeared essentially identical on both selective and nonselective media. All callus from seven plates of the

pHYGI1/pBII221 and one T.E. treated plate were transferred from round 1 selection plates to round 2 selection plates that contained 60 mg/l hygromycin. The round 2 selection plates each contained ten 30 mg pieces of callus per plate, resulting in an expansion of the 35 total number of plates.

After 21 days on the round 2 selection plates, all of the material was transferred to round 3 selection

plates containing 60 mg/l hygromycin. After 79 days post-bombardment, the round 3 sets of selection plates were checked for viable sectors of callus. One of the sectors was proliferating from a background of necrotic tissue on plates treated with pHYGI1/pBII221. The sector was designated PH3 and was transferred to F medium without hygromycin.

After 19 days on F medium without hygromycin, PH3 was transferred to F-medium containing 60 mg/l 10 hygromycin. PH3 was found to be capable of sustained growth through multiple subcultures in the presence of 60 mg/l hygromycin.

V. Confirmation of transformed callus

15 To show that the PH3 callus had acquired the hygromycin resistance gene, genomic DNA was isolated from PH3 callus and unselected control callus in accord with Example V, and analyzed by Southern blotting. The isolated DNA (10 μg) was digested with BamHI (NEB) and 20 electrophoresed in a 0.8% w/v agarose gel at 15 V for 16 hrs in TAE buffer (40 mM Tris-acetate, pH 7.6, 1 mM EDTA). The DNA was transferred to a Nytran membrane (Schleicher and Schuell). Transfer, hybridization and washing conditions were carried out as per the manufacturer's recommendations.

A 32 P labelled probe was prepared by random primer labelling with an Oligo Labelling Kit (Pharmacia) as per the supplier's instructions with $\alpha-^{32}$ P-dCTP (ICN Radiochemicals). The template DNA used was the 1055 bp BamHI fragment of pHYGI1, which contains the entire HPT coding sequence.

Membranes were exposed to Kodak X-OMAT AR film in an X-OMATIC cassette with intensifying screens. A band was observed for PH3 callus at the expected

35 position of 1.05 Kb, indicating that the HPT coding sequence was present. No band was observed for control callus.

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To demonstrate that the hygromycin gene is incorporated into high molecular weight DNA, undigested DNA from PH3 callus and control callus was electrophoresed, blotted and hybridized as described 5 above. Undigested PH3 DNA only showed hybridization to the probe at a mobility equal to uncut DNA. No hybridization was observed for DNA from control callus. These results demonstrate that the HPT coding sequence is not present in PH3 callus as intact pHYGI1 or as a small non-chromosomal plasmid. The results are consistent with incorporation of the hygromycin gene into high molecular weight DNA.

VI. Plant regeneration and production of seed

Portions of PH3 callus were transferred directly from plates containing 60 mg/l hygromycin to RM5 medium which consists of MS basal salts (Murashige et al. 1962) supplemented with thiamine HCl 0.5 mg/l, 2,4-D 0.75 mg/l, sucrose 50 g/l, asparagine 150 mg/l, and Gelrite 2.5 g/l (Kelco Inc., San Diego).

After 14 days on RM5 medium, the majority of PH3 and unselected control callus was transferred to R5 medium (RM5 medium, except that 2,4-D is omitted). The plates were cultured in the dark for 7 days at 26°C and transferred to a light regime of 14 hrs light and 10 hrs dark for 14 days at 26°C. At this point, plantlets that had formed were transferred to one quart canning jars (Ball) containing 100 ml of R5 medium. Plants were transferred from jars to vermiculite for 7 or 8 days before transplanting them into soil and growing them to maturity. A total of 45 plants were produced from PH3 and a total of 10 plants were produced from control callus.

Controlled pollinations of mature PH3 plants

35 were conducted by standard techniques with inbred Zea

mays lines MBS501 (Mike Brayton Seeds), FR4326 (Illinois
Foundation Research) and the proprietary inbred line

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LM1112. Seed was harvested 45 days post-pollination and allowed to dry further for 1-2 weeks.

VII. Analysis of the R1 progeny

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R1 plants were tested for the presence of the HPT and GUS gene sequences by PCR analysis. Expression of the HPT gene was determined with an enzymatic assay for HPT activity. Table 3 summarizes the data. These data demonstrate transmission and expression of the 10 recombinant DNA through both male and female parents, and are consistent with Mendelian inheritance. Southern blot evidence for integration of the HPT coding sequence into high molecular weight DNA for the source callus, combined with the inheritance data, suggest that 15 the foregoing DNA sequences are chromosomally integrated. The presence of GUS gene sequences in the R1 progeny demonstrates cotransformation and inheritance of an unselected gene.

Table 3 Analysis of R1 Progeny of PH3

Transformants

Parents: MBS501 X PH3.4

	Enzyme	PCR	Assay
PH3.4 Plant	<u>HPT Assay</u>	<u>HPT</u>	<u>GUS</u>
4 1			
4.1	-	_	_
4.2	+	+	+
4.3	+	+	+
4.4	-	-	-
4.5	+	+	+
4.6	+	+	+
4.7	+	+	+
4.8	-	-	-
4.9	-	-	-

	Parents:	LM1112	X P	H3.18	
18.1	-		-	•	_
18.2	+		4	-	+
18.3	_		_	•	_
18.4	_		_	•	_
18.5	+		4	•	+
18.6	+		4	_	<u>.</u>
18.7	+		4	•	+
18.8	+		4	_	÷
18.9	+		د.	_	
18 10				_	T .
15.11			1		

Parents: PH3.2 X FR4326

_	Enzyme	PCR	Assay
<u>Plant</u>	HPT Assay	HPT	GUS
2.1	-	_	_
2.2	-	-	_
2.3	+	+	+
2.4	_	_	_
2.5	+	+	+
2.6		-	-
2.7	+	+	+
2.8	+	+	ND
2.9	+	+	ND

<u>Controls</u>

Parents: TE3.1 X Oh43

_			
3.1	-	-	ND
3.2	-	-	ND
3.3	-	_	ND
3.4	_	_	ND
3.5	_	_	ND
3.6	_	_	
3.7	_	_	ND
	_	_	-
3.8	-	_	-
3.9	-	_	_
3.10	-	· -	_
3.11	-	ND	ND
3.12	_	ND	ND
3.13		ND	ND
3.14	-	ND	ND
3.15	_	ND	ND
3.16			
	-	ND	ND
3.17	-	ND	ND
3.18	-	ND	ND
3.19	-	ND	ND
		. –	

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3.20	-	ND	ND
3.21	-	ND	ND
3.22	-	ND	ND
3.23	-	ND	ND
3.24	-	ND	ND

The HPT enzyme assay was based on the following methods: S. K. Datta et al., Bio/Technology, 8, 736 (1990), M. Staebell et al., Anal. Biochem., 185, 319 (1990), and E. Cabanes-Bastos, Gene, 77, 169 (1989).

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Root samples (0.2 g) were excised from 7-10 day old seedlings and were quick frozen in liquid nitrogen . (N_2). Samples were ground with alumina and 400 μ l of EB (50mM Tris-HCl, 10% v/v glycerol, 0.1 mM PMSF, pH 7.0) for 30-60 seconds using a disposable pestle and tube 10 (Kontes), then centrifuged in an Eppendorf microfuge for 10 min. The supernatant was transferred to a Centricon 30 filter unit and desalted by centrifuging in a fixed angle rotor at 5000 rpm in a Beckman GPR centrifuge for 30 min at 4°C. One ml of EB was used to wash each 15 filter unit, spinning an additional 5000 RPM for 60 min. The retentate was then recovered and stored at -70°C. Samples of transgenic and unselected control callus tissue (0.2 g) were ground as described above, and the supernatents were used as positive and negative 20 controls.

Protein was quantified using the method of Bradford, Anal. Biochem., 72, 248 (1976) with a BioRad kit. Protein concentrations in root extracts ranged from $0.2-2 \, \mu g/\mu l$.

25 The root extract (4 µg total protein) was added to a reaction mixture containing 20 mM Tris-maleate, 13 mM MgCl₂, 120 mM NH₄Cl, 0.5 mM DTT, 50 μ M ATP, 0.61 μ g/ μ l hygromycin B, 25 $\mu g/\mu l$ bovine serum albumin, and 12 μCi gamma 32P-ATP. Reaction volume was 33 μ l. The reaction 30 mixture was incubated for 30 min at 37°C.

One µl of the reaction mixture was spotted on a polyethyleneimine cellulose thin layer chromatography

plate (Sigma Chem. Co.). Plates were developed in 50 mM formic acid pH 5.4, air dried, and exposed to Kodak XAR-5 film. The reaction product, hygromycin phosphate, migrates near the solvent front under these conditions.

To conduct the PCR assay, 0.1g samples were taken from plant tissues and frozen in liquid nitrogen. Samples were then ground with 120 grit carborundum in 200µl 0.1M Tris-HCl, 0.1M NaCl, 20mM EDTA, 1% Sarkosyl pH 8.5) at 4°C. Following phenol/chloroform extraction and ethanol and isopropanol precipitations, samples were suspended in T.E. and analyzed by polymerase chain reaction (K.B. Mullis, U.S. Pat. No. 4,683,202).

PCR was carried out in 100 µl volumes in 50 mM KCl, 10 mM Tris-HCl pH 8.4, 3 mM MgCl₂, 100 µg/ml

15 gelatin, 0.25 µM each of the appropriate primers, 0.2 mM of each deoxynucleoside triphosphate (dATP, dCTP, dGTP, dTTP), 2.5 Units of Taq DNA polymerase (Cetus), and 10 µl of the DNA preparation. The mixture was overlaid with mineral oil, heated to 94°C for 3 min, and

20 amplified for 35 cycles of 55°C for 1 min, 72°C for 1 min, 94°C for 1 min. The mixture was then incubated at 50°C for 2 min and 72°C for 5 min. 10 µl of the PCR product was electrophoresed in agarose gels and visualized by staining with ethidium bromide.

PCR primer complementary to the 35S promoter, and one complementary to the HPT coding sequence were employed. Thus, in order to generate the appropriately sized PCR product, the HPT template DNA must contain contiguous 35S promoter region, Adhl intron, and 5' protein coding sequence region.

PCR primers complementary to sequences within the GUS protein coding region were employed. A 797 bp PCR product is predicted if the template DNA contains an intact coding region between the two primers.

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EXAMPLE II

Fertile Transgenic Plants from Callus Line AB63S Containing Recombinant DNA Encoding a Seed Storage Protein

The 10 kD-zein storage protein is produced in the endosperm of the maize kernel and is a representative member of a class of proteins referred to as "seed storage proteins." The protein contains extremely high levels of methionine (22.5%) and is 10 encoded by the Zps10/(22) gene (M.S. Benner et al., Theor. Appl. Genet., 78, 761 (1989)); a gene referred to as z10 herein. Thus, increased expression of the z10 gene can be used to increase the methionine content of corn. Production of fertile transgenic corn containing 15 a chimeric z10 gene was accomplished by the procedures of Example I with some minor modifications. Example II also demonstrates the introduction of recombinant DNA using a callus line different from that used in Example I.

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I. Tissue Culture Lines

The embryogenic maize callus line named AB63S used in this example was produced from immature embryos resulting from a cross between elite inbred lines A188 25 and B73, by the initiation and maintenance procedures described in Example I. The callus line had been initiated approximately 7 months prior to bombardment. Eleven weeks prior to bombardment the tissue was sieved by forcing it through a 1.9 mm screen and plated onto N6 maintenance medium. Friable, embryogenic callus was selected from the resulting tissue after 1-2 weeks of growth, transferred to fresh medium and allowed to grow for 1.5-3 weeks before sieving again. This cycle of sieving and recovery of friable callus was carried out a 35 total of three times prior to subjecting the tissue to bombardment.

II. Plasmids

The plasmids pHYGI1 and pBII221 described in Example I were used in this Example. In addition, the plasmid pZ27Z10 was included in the DNA/tungsten 5 preparations. The plasmid pZ27Z10 was constructed using the vector puclis (J. Vieira et al., Methods Enzymol., 153, 3 (1987), a 3.2 kb circular plasmid, by standard recombinant DNA techniques. pZ27Z10 contains the 5' transcriptional regulatory region from a maize 10 27 kD-zein gene (D.E. Geraghty, Ph.D. Thesis, University of Minnesota (1987)), referred to as z27 in this document), positioned immediately adjacent to the coding sequence and 3' noncoding sequence from a maize 10 kD-zein gene (Kirihara et al., <u>Gene</u>, <u>7</u>, 359 (1988)), The combination referred to as z10 in this document). of the z27 regulatory sequence and z10 coding and 3' sequences is referred to as the chimeric z27-z10 gene in this document. The poly A signal from the region of the CaMV genome adjacent to the 35S promoter is positioned 20 3' to the z10 gene sequences in pZ27Z10. Figure 7 depicts a map of this plasmid.

A sample of plasmid pZ27Z10 has been deposited in the American Type Culture Collection, Rockville, MD, under the provisions of the Budapest Treaty under 25 accession number ATCC 40938.

III. DNA Delivery Process

Tissue from the maize embryogenic callus line
AB63S was subcultured 3 weeks prior to microprojectile
30 bombardment. The tissue was prepared for bombardment by
sieving it through a 1.9 mm screen. The sieved tissue
was plated onto sterile 5.5 cm Whatman No. 1 filter
disks as a thin lawn of approximately 500 mg of tissue
per filter disk. A total of 6 disks of tissue were
35 prepared. Prior to microprojectile bombardment the
filter disks containing the tissue were transferred to
empty petri plates. The tissue was dehydrated slightly

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by allowing the plates to stand uncovered in a laminar flow hood for 20 minutes. To prevent further dehydration of the tissue, the plates were stored in a high humidity box until used for bombardment.

The DNA used in the bombardment consisted of a 1:1:1 mixture (by weight) of the plasmids pHYGI1, pZ27Z10 and pBII221. Precipitation of the DNA onto tungsten particles was carried out as described in Example I.

in Example I except that the Biolistic™ PDS 1000
(DuPont) instrument was used, and no screen was placed over the tissue. The six filter disks with callus tissue were treated as follows: two filters were
bombarded two times each with the DNA/tungsten preparation (potential positive treatment 2X); three filters were bombarded one time each with the DNA/tungsten preparation (potential positive treatment 1X); one filter was bombarded once with a tungsten/water suspension using the same weight of tungsten used in the DNA/tungsten bombardments (negative control treatment).

IV. Selection of Transformed Callus

25 tissue from the 2X DNA treatment were transferred to round 1 selection plates: F medium containing 15 mg/l hygromycin. Two of the filters from the 1X DNA treatment and the filter from the negative control treatment were also transferred to round 1 selection 30 medium. The third filter from the 1X DNA treatment was transferred to F medium containing no hygromycin, for use as unselected control callus. Since this unselected control callus was potentially positive, it was only maintained for the first two rounds of selection for comparative purposes; subsequently, non-bombarded AB63S callus maintained on F medium without hygromycin was used as unselected control callus.

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After five days the callus was transferred in small clumps (about 25 mg) from the filter disks onto fresh round 1 selection medium. The plating density was 10 callus clumps per plate. By this time the callus had 5 increased approximately two-fold in weight. unselected control callus was transferred in the same manner to F medium without hygromycin. At nineteen days post-bombardment, all of the tissue was transferred from round 1 selection medium to round 2 selection 10 medium (F medium containing 60 mg/l hygromycin). plating density was 14 callus clumps per plate at this transfer. The callus had increased approximately three-fold in volume during the 14 days since the previous transfer. After 23 days, all of the callus was 15 transferred from round 2 selection medium to round 3 selection medium containing 60 mg/l hygromycin.

At 59 days post-bombardment, five live sectors were observed proliferating among the surrounding necrotic callus clumps. These five sectors were thought to have been derived from a single callus clump from the round 2 selection, since they appeared adjacent to one another on consecutive plates from the previous transfer. These sectors arose from a 2X DNA treatment and were designated as callus line Met1.

The callus line Met1 was transferred to F
medium containing 60 mg/l hygromycin and to F medium
without hygromycin. After 22 days of incubation, there
was no visible difference in growth or appearance of the
tissue on the two types of media. The callus line Met1
grew rapidly and appeared to be uninhibited by the
presence of the hygromycin in the medium. The Met1
callus appeared highly friable and embryogenic on either
medium, and was visually indistinguishable from control
AB63S callus grown on F medium without hygromycin.

V. Confirmation of Transformed Callus

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An inhibition study was performed using Metl

callus and control AB63S callus. Prior to initiating the inhibition study, Metl callus was grown for 21 days on F medium without hygromycin. Callus of Metl and control AB63S callus was transferred onto plates of F medium containing 0, 15, 60, 100 and 200 mg/l hygromycin. Three plates of each callus line were prepared for each concentration of hygromycin; each plate contained 5 or 6 pieces of callus at approximately 50 mg per piece (total callus per plate was approximately 300 mg).

After 28 days of incubation, the weight of callus on each plate was measured. A measure of percent growth inhibition was determined by dividing the average weight of callus obtained at each hygromycin

15 concentration by the average weight of callus obtained at 0 mg/l hygromycin. The results showed that the growth of Metl callus was completely uninhibited at hygromycin concentrations of 15, 60 and 100 mg/l. The growth of Metl callus was approximately 20% inhibited on 20 medium containing 200 mg/l hygromycin. In contrast, the growth of AB63S callus was inhibited at all concentrations of hygromycin; at 200 mg/l hygromycin, AB63S callus was 90% inhibited in growth. These results confirmed that the callus line Metl exhibited resistance to the presence of hygromycin in the growth medium.

Southern blot analysis was carried out to verify the presence of the hygromycin resistance gene in Metl callus DNA and to determine whether the callus had also acquired the chimeric z27-z10 gene. For detection of the HPT coding sequence, DNA samples isolated from Metl callus and from control callus were digested with the restriction enzymes BamHI, HindIII and BstEII. The digested DNA samples were subjected to electrophoresis on a 0.8% agarose gel and blotted to a Nytran membrane as described in Example I. Visual inspection of the ethidium bromide-stained gel prior to blotting indicated that the BamHI digestions appeared to be complete,

whereas neither HindIII nor BstEII cleaved the DNA samples to a significant degree. The blot was probed with a biotin-labeled 1.05 kb BamHI fragment from the HPT coding sequence. The conditions used for 5 hybridization, washing and detection were as suggested in a BRL Photogene™ kit used in this experiment.

After hybridization, labeled bands were observed in the lanes containing BamHI digested Metl DNA at the expected size of 1.05 kb as well as at sizes of 10 approximately 2.7, 4.5 and 6.5 kb. This result showed that the HPT coding sequence was present in DNA from the callus line Met1. The additional bands at 2.7, 4.5 and 6.5 kb indicated that either the restriction enzyme digestion was incomplete, or that multiple rearranged 15 copies of the HPT sequence were present. For the HindIII and BstEII digests, hybridization signals were visible in both lanes at the position of undigested DNA. This result indicated that the HPT coding sequence was integrated into high molecularweight DNA in the Metl 20 genome. No hybridization signals were observed in any of the lanes containing digestions of DNA from control callus.

For detection of the chimeric z27-z10 gene,
Southern blot analysis was carried out using DNA samples
isolated from Metl callus and control callus. The DNA
samples were digested with the restriction enzymes BamHI
and EcoRI. A BamHI digestion liberates a 2.76 kb
fragment containing the z10 coding sequence and 3'
noncoding sequence from the pZ27Z10 construction used in
this transformation. There is a single EcoRI site in
the 7.26 kb pZ27Z10 plasmid. The blot prepared was
hybridized with a biotin-labeled probe containing the
entire z10 coding sequence, using the conditions
described in a Photogene kit (BRL).

35 For the BamHI digestions, endogenous z10 sequences gave hybridization signals at 9 kb and at approximately 15 kb in the lanes containing both Met1

and control callus DNAs. An additional band was observed at approximately 3.5 kb in the Met1 sample but not in the control DNA samples. No strong hybridization signal was observed at the expected 2.76 kb size in Met1 DNA. The lack of a labeled band of 2.76 kb in the BamHI digestion of Met1 DNA indicated either incomplete digestion of the DNA, or rearrangement of the introduced DNA.

For the EcoRI digestions, endogenous z10

10 sequences gave hybridization signals at approximately 12 kb and 16 kb in both Metl and control callus DNAs. An additional band was observed at approximately 14 kb in the Metl sample but not in the control callus DNA samples. These results indicated that novel z10

15 sequences were present in DNA from Metl callus that were absent from control callus DNA samples.

These results were confirmed by a second Southern blot analysis in which Metl and control callus DNAs were digested with BamHI, NcoI and NsiI. Also 20 included were samples of undigested DNA from Metl and control callus. The filter was probed with a 32P-labeled z10 coding sequence probe. In all of the digestions, novel z10 hybridization signals were observed in Met1 callus DNA that were absent in the negative control 25 callus DNA samples. These results confirmed that novel z10 coding sequences were present in the Met1 callus. In the lanes containing undigested DNA, hybridization signals were observed at the position of undigested DNA These results from both Metl and control callus. 30 indicated that the introduced z10 sequences were integrated into chromosomal DNA in Metl callus.

VI. Plant Regeneration and Production of Transgenic Seed

Callus from the line Met1 and from the control line AB63S was placed on RM5 medium as described in Example I. After 14 days of incubation at 26°C in the

dark, the callus was transferred to R5 medium as in Example I. The callus was incubated at 26°C in the dark for 7 days, and then transferred to the light under the conditions described in Example I. After 14 days in the light, plantlets were transferred to Magenta boxes (Sigma Chemical Co.) containing 100 ml of R5 medium. After 7-14 days on this medium, plantlets were transferred to vermiculite for 7 days and then to soil where they were grown to maturity. A total of 49 plants (designated Met1-1 through Met1-49) were regenerated from Met1 callus, and a total of 25 plants were regenerated from control AB63S callus.

PCR analysis was carried out on samples of DNA prepared from leaf tissue to verify that the plants

15 regenerated from Metl callus had retained the introduced DNA. For this purpose, a set of six oligodeoxyribonucleotides were used. The sequences of these oligonucleotides, their orientations and relative positions within the chimeric z27-z10 gene construction shown in Figure 8 are summarized in Table 4 below.

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Table 4

I. OLIGONUCLEOTIDE NAMES/POSITIONS IN CHIMERIC Z27-Z10 GENE:

1.	Z27-5'	nt	132-156
2.	Z27-MID	nt	628-651
3.	Z27-3'	nt	1081-1104
4.	Z10-5'	nt	1182-1206
5.	Z10-MID	nt	1420-1444
<u> </u>	710-31	n.t	1548-1571

II. AMPLIFIED FRAGMENT SIZES FROM OLIGONUCLEOTIDE PAIRS:

A. ENDOGENOUS OR CHIMERIC GENE PAIRS:

(1)	+	(3)	972	BF
(2)			476	BP
(4)			389	BF
(4)			262	BF

B. CHIMERIC GENE-SPECIFIC PAIRS:

(1)	+	(6)	1439	BP
(1)			1312	BP
(2)			943	BP
(2)			816	BP

The set of oligonucleotides (or primers) was designed such that the use of a pair of oligonucleotides consisting of one z27 oligonucleotide (z275' or z27mid) and one z10 oligonucleotide (z10mid or z103') would 5 result in amplification of fragments from the chimeric z27-z10 gene only, and not from endogenous maize z27 or z10 genes. The appearance of amplified fragments of the expected sizes in PCR reactions using these z27-z10-specific oligonucleotide pairs were diagnostic for the presence of the chimeric z27-z10 gene in a particular callus or plant DNA sample.

The use of a pair of z27 oligonucleotides (z275' or z27mid with z273') or a pair of z10 $\,$

oligonucleotides (z105' with z10mid or z103') resulted in amplification of fragments representing z27 regulatory sequences or z10 coding sequences, respectively, when used in PCR reactions carried out with control AB63S callus DNA, Met1 callus DNA or pz27z10 plasmid DNA. These reactions served as positive controls for amplification of endogenous maize gene sequences from a particular callus or plant DNA sample.

Leaf DNA was prepared from two Met1 R0 plants 10 (designated Met1-1 and Met1-2) and from one control AB63S RO plant as described hereinbelow. PCR reactions were carried out with these DNA samples using the oligonucleotide pairs specific for the HPT coding sequence, the chimeric z27-z10 gene, the z27 regulatory 15 sequence and the z10 coding sequence. Gel analysis of the PCR reaction products showed that the expected size amplification products were obtained in all reactions using Met1-1 DNA and Met1-2 DNA. Control AB63S DNA was negative for the HPT coding sequence and for the 20 chimeric z27-z10 gene, but positive for the endogenous z27 regulatory sequence and z10 coding sequence. results demonstrated that the Metl RO plants examined had retained the introduced HPT DNA and the chimeric z27-z10 DNA.

25 As corroborative proof that the diagnostic z27-z10 PCR reaction products contained both z27 regulatory sequences and z10 coding sequences, Southern blot analysis was carried out on the PCR reaction products. Southern blot filters were prepared from two agarose gels loaded with identical samples. The samples contained the PCR reaction products resulting from amplification of z27 regulatory sequences, z10 coding sequences, and chimeric z27-z10 gene sequences described above. One blot was hybridized with a z10 coding sequence probe and the other blot was hybridized with a z27 regulatory sequence probe. The results showed that the z10 coding sequence probe hybridized to the

amplified fragments from the z10 coding sequence PCR reactions and the chimeric z27-z10 gene PCR reactions, but not to the amplified fragments from the z27 regulatory sequence PCR reactions. Analogously, the z27 probe hybridized to the amplified fragments from the z27 regulatory sequence PCR reactions and the chimeric z27-z10 gene PCR reactions but not to the amplified fragments from the z10 coding sequence PCR reactions. These results demonstrated that the fragments amplified in the PCR reactions contained the expected gene sequences and that the diagnostic PCR amplification product for the chimeric z27-z10 gene contained both z27 regulatory sequences and z10 coding sequences.

Controlled pollinations of mature Met1 R0

15 plants were carried out with inbred maize lines A188,
B73, A654 and H99. In addition, several self- and
sib-pollinations were carried out. A total of 130
pollinations were carried out, using Met1 R0 plants as
both male pollen donors and as female pollen recipients.

20 Mature seed was harvested at 45 days post-pollination
and allowed to dry further for 1-2 weeks.

VII. Analysis of the R1 progeny

The presence of the HPT sequence and the
chimeric z27-z10 gene were evaluated by PCR analysis of
DNA from R1 plant tissues, in three sets of R1 progeny.

The first set of R1 progeny analyzed consisted of four immature tassel-seeds from R0 plant Met1-7.

These tassel-seeds were obtained through open

30 pollination of silks developing from the tassel of Met1-7. Approximately 18 days after pollination, the tassel-seeds were removed from the plant and surface-sterilized. The endosperm from each seed was excised and frozen on dry ice and stored at -70°C until used for DNA isolation.

The embryo from each seed was transferred to R5 medium and allowed to germinate. After 10 days, the

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seedlings were transferred to vermiculite for 7 days and then to soil, where they were allowed to grow to maturity. PCR analysis was carried out on DNA isolated from each endosperm using oligonucleotide sets specific 5 for the HPT coding sequence, the chimeric z27-z10 gene and the maize Adh-1 gene. Three of the four endosperm DNA samples were found to be positive for all of the above gene sequences. The fourth endosperm DNA sample was negative for the HPT coding sequence and the chimeric z27-z10 gene sequence but positive for the endogenous Adh-1 sequence. These results showed that both the HPT sequence and the chimeric z27-z10 gene had been transmitted to R1 progeny of Met1-7.

A similar analysis was carried out with 24

15 seeds from the cross A654 X Met1-1. At 24 days after pollination, the seeds were removed from the ear and surfacesterilized. Endosperms and embryos were isolated and treated as described above. PCR analysis of the endosperm DNA samples showed that 9 of the 24 progeny

20 had received both the HPT sequences and the chimeric z27-z10 gene. The remaining 15 progeny carried neither of the introduced gene sequences. The third set of R1 progeny analyzed consisted of 28 R1 plants derived from self-pollination of R0 plant Met1-6. DNA samples were

25 prepared from leaf tissue of 28 two-week old seedlings.

PCR analysis of the leaf DNA samples showed that 24 of the R1 progeny carried both the HPT sequences and the chimeric z27-z10 gene. The remaining 4 R1 progeny carried neither of the introduced gene

30 sequences. The results of the analyses described above are summarized in Table 5 and demonstrate that a significant proportion of the R1 progeny of Metl plants inherited the introduced DNA.

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Table 5

Inheritance of the Introduced HPT Sequences and the Chimeric Z10 Gene in Progeny from Crosses of Metl Plants

		Outcross (<u>A654 X Met1-1</u>)	Self-Pollination (Met1-6)
10	<u>Genotype</u>	No. Progeny	No. Progeny
15	HPT+/z27z10+ HPT+/z27z10- HPT-/z27z10+ HPT-/z27z10-	9/24 0/24 0/24 15/24	24/28 0/28 0/28 4/28

The ratio of z27z10+/HPT+ plants to z27z10
20 /HPT- plants from the outcross population is consistent with a 1:1 segregation ratio of a single locus. In addition, the ratio of z27z10+/HPT+ plants to z27z10-/HPT- plants from the self-pollinated population is consistent with a 3:1 segregation ratio of a single

25 locus. The data also indicate linkage of the introduced HPT and z27-z10 sequences.

EXAMPLE III

Fertile Transgenic Plants from Callus Line AB12 Containing Recombinant DNA Encoding an Insecticidal Protein

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The proteins encoded by the various Bacillus thuringiensis genes have been shown to be useful in a number of pesticidal applications. Production of transgenic maize plants with specific heterologous Bt genes may improve the resistance of the plants to specific pests. Fertile transgenic plants containing a recombinant Bt gene were prepared according to the procedure of Example I, with several minor modifications, as indicated below.

At the same time AB11, the callus line of Example I, was initiated, another callus line was also prepared. AB12 was prepared from a separate cross of plants with the same parental genotype (A188 X B73).

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This example demonstrates the regeneration of fertile transgenic plants from a third callus line in addition to AB11 and AB63S, and demonstrates that the production of fertile transgenic plants was not the result of properties unique to a single callus line.

The plasmid pHYGI1 described in Example I was used, as well as pMS533, which is a plasmid that contains the insecticidal <u>Bacillus thuringiensis</u> endotoxin (BT) gene fused in frame with the neomycin phosphotransferase (NPTII) gene. At a position 5' from the fusion gene are located segments of DNA from the CaMV and nopaline synthase promoters. At a position 3' from the fusion gene are segments of DNA derived from the tomato protease inhibitor I gene and the poly A region of the nopaline synthase gene.

Callus was bombarded as described in Example I except that the DNA used in the tungsten/DNA preparations contained plasmids pHYGI1 and pMS533. One tube contained only 5 μ l TE (10 mM Tris-HCl pH 8.0, 1 mM EDTA).

The following bombardments were done: 11 \times pHYGI1/pMS533 (potential positive treatment) and 3 \times TE prep (control treatment).

After bombardment, the callus from the pHYGI1/

pMS533 treatments was placed onto round 1 selection
plates, F medium containing 15 mg/l hygromycin as ten
pieces per plate. The same was done for two of the
plates bombarded with the TE preparation (selected
control callus). One plate of callus bombarded with the

TE preparation was placed onto F medium with no
hygromycin; this callus was maintained throughout the
experiment as a source of control tissue (unselected
control callus).

After 18 days, the callus was transferred from 5 round 1 selection plates to round 2 selection plates containing 60 mg/l hygromycin as ten 30 mg pieces per plate. After 21 days of selection on round 2 selection

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plates, the callus appeared completely inhibited. All of the callus was transferred from round 2 selection plates to round 3 selection plates containing 60 mg/l hygromycin.

After 42 days on round three selection plates, six sectors were observed proliferating from the surrounding necrotic tissue, all were from pHYGI1/pMS533-treated material. The callus lines were transferred to F medium.

After 24 days, one callus line, designated CB2, had grown substantially. Portions of each callus line were then transferred to (i) F medium containing 15 mg/l hygromycin, or (ii) F medium containing 60 mg/l hygromycin. Control callus was plated on F medium with 15 15 mg/l hygromycin.

Only one of the six callus lines, CB2, was capable of sustained growth through multiple subcultures in the presence of 60 mg/l hygromycin. DNA was isolated from portions of this callus and the presence of the Bt 20 gene was confirmed by Southern blot analysis.

DNA was digested with the restriction enzymes BamHI and XhoI in combination. BamHI cuts at the 5' end of the Bt coding sequence and XhoI cuts at the 3' end of the coding region. A 32P labeled probe was prepared from 25 the 1.8kb BamHI/XhoI restriction fragment of pMS533. The predicted 1.8 kb band was observed after hybridization and autoradiography. No bands were observed in the DNA from the untransformed callus.

Plants were regenerated from CB2 and controlled 30 pollinations were performed to yield the R1 generation, as described hereinabove. R1 plants were tested for the presence of the HPT and Bt gene sequences by PCR analysis, and for HPT gene expression by enzyme assay as described in Example I.

35 PCR analysis was performed on DNA isolated from root tissue of R1 seedlings by the methods described in Example I. To test that each DNA sample was competent

for PCR, PCR analysis was carried out using primers specific for the native maize 10kD zein gene. The primers used for PCR analysis of the Bt gene were one complementary to the 35S promoter and one complementary to the Bt coding sequence. Primers used for HPT PCR analysis were the same as those described in Example I. The results of this analysis are presented in Table 6. The results show transmission of the introduced HPT and Bt genes through both the male and female parents and the observed frequencies are consistent with Mendelian inheritance. The results are consistent with the insertion of the HPT and Bt genes into chromosomal DNA. It was also found that the HPT and Bt genes show a linked inheritance, indicating that the two genes are inserted close together on the same chromosome.

Table 6

Analysis of R1 Progeny of CB2

A. Transgenic CB2.8 x FR4326				Control AB12.7 x FR4326					
	Ana	lysi	<u>s</u>			2	Analy	sis	
<u>Plant</u>	10Kd PCR		BT PCR	HPT ENZ.	PLANT	10Kd PCR	HPT PCR	BT PCR	HPT ENZ.
1 2 3 4 5 6 7 8 9	+ + + + + + + + + + + + + + + + + + + +	+ + + - + + - + + - + - + + +	+ + + + + + + + + + + + + + + + + + + +	+ + - + + - + -	1 2	+ +	-	-	-

	B. Transgenic CB2.11 x LB101				Control AB12.4 x LB101					
	<u>Assa</u>	<u>ys</u>				Assa	ays			
Plant	10Kd PCR		BT PCR	HPT ENZ.	PLANT		HPT PCR		HPT ENZ.	
1 2 3 4 5 6 7 8 9 10	+ + + + + + + + +	+ - +	- +	_	1 2	++	-	-		
	C. Transgenic CB2.7 x LM1112 Assays					Control <u>AB12.5 x LM1112</u> <u>Assays</u>				
<u>Plant</u>	10Kd PCR	HPT PCR			<u>PLANT</u>	10Kd PCR	HPT PCR	BT PCR		
1 2 3 4 5 6 7 8 9		- +	++		1 2	+	Ξ	-		

D.	Transgenic	
	FR4326 x CB2.	9

Control FR4326 x AB12.1

	<u>Assays</u>			<u>Assays</u>				
<u>Plant</u>	10Kd PCR		BT PCR		PLANT.	10Kd PCR	HPT PCR	BT PCR
1 2 3 4 5 6 7 8 9	+ + + + + + + +	- + + + + - -	- + + + + + - +		1 2	+ '	Ξ	-

E. Transgenic A188 x CB2.1

Control A188 x AB12.2

Assays

<u>Assays</u>

Plant	10Kd PCR			PLANT	10Kd PCR	HPT PCR	BT PCR
1	+	+	+	1	+	_	_
2	+	+	+	2	+	_	_
3	+	+	+				
4	+	_	-				
5	+	+	+				
6	+	_	_				
7	+	+	+				
8	+	+	+				
9	+	_	_				
10	+	-	_				

F. Tra	Control <u>LM112 x AB12.6</u>						
	Assays			<u>Assays</u>			
Plant	10Kd PCR			PLANT	10Kd PCR	HPT PCR	BT PCR
1 2 3 4 5 6 7 8	+ + + + + + + + +	- - - - - +	- - - - - - +	1 2	+ +	-	-

EXAMPLE IV

Fertile Transgenic Plants from Callus Line AB12 Transmitting Recombinant DNA through Two Complete Sexual Cycles

5 I. Plant Lines and Tissue Culture

The callus line AB12 was used, and is described in Example III.

II. Plasmids

The plasmids pBII221 and pHYGI1 described in Example I were used.

III. DNA Delivery Process

Callus was bombarded as in Example I, except

that the DNA used in the tungsten/DNA preparations differed. All of the tubes contained 25 µl 50 mg/ml M10 tungsten in water, 25 µl 2.5 M CaCl₂, and 10 µl 100 mM spermidine along with a total of 5 µl 1 mg/ml total plasmid content. One tube contained only plasmid

20 pBII221; two tubes contained only plasmid pHYGI1; and one tube contained no plasmid but 5 µl TE buffer.

The following bombardments were done: 2 x pBII221 prep (for transient expression); 7 x pHYGI1 prep

(potential positive treatment); and $3 \times TE$ prep (negative control treatment).

After all the bombardments were performed, the callus from the pBII221 treatment was transferred plate for plate to F medium as five 50 mg pieces. After 2 days, the callus was placed into GUS assay buffer as per Example I. Numbers of transiently expressing cells were counted and found to be 686 and 845 GUS positive cells, suggesting that the particle delivery process had occurred in the other bombarded plates.

IV. Selection of Transformed Callus

After bombardment, the callus from the pHYGI1 treatments was placed onto round 1 selection plates,

F medium containing 15 mg/l hygromycin, as ten 25 mg pieces per plate. The same was done for two of the plates bombarded with the TE preparation (selected control callus). One plate of callus bombarded with the TE preparation was placed onto F medium with no hygromycin; this callus was maintained throughout the experiment as a source of control tissue (unselected control callus).

After 13 days, the callus on round 1 selection plates was indistinguishable from unselected control callus. All of the callus was transferred from round 1 selection plates to round 2 selection plates containing 60 mg/l hygromycin. An approximate five-fold expansion of the numbers of plates occurred.

The callus on round 2 selection plates had increased substantially in weight after 23 days, but at this time appeared close to necrotic. All of the callus was transferred from round 2 selection plates to round 3 selection plates containing 60 mg/l hygromycin.

At 58 days post-bombardment, three live sectors were observed proliferating from the surrounding dead tissue. All three lines were from pHYGI1 treatments and were designated 24C, 56A, and 55A.

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After 15 days on maintenance medium, growth of the lines was observed. The line 24C grew well whereas lines 55A and 56A grew more slowly. All three lines were transferred to F medium containing 60 mg/l 5 hygromycin. Unselected control callus from maintenance medium was also plated to F medium containing 60 mg/l hygromycin.

After 19 days on 60 mg/l hygromycin, the growth of line 24C appeared to be entirely uninhibited, while 10 the control showed approximately 80% of the weight gain The line 56A was completely necrotic, and the line 55A was very close to necrotic. The lines 24C and 55A were transferred again to F medium containing 60 mg/l hygromycin, as was the control tissue.

After 23 days on 60 mg/l hygromycin, the line 24C again appeared entirely uninhibited. The line 55A was completely dead, as was the negative control callus on its second exposure to F medium having 60 mg/l hygromycin.

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V. Confirmation of Transformed Callus

A Southern blot was prepared from BamHIdigested DNA from the line 24C and probed with the HPT probe described in Example I. As shown in Figure 6, a 25 band was observed for the line 24C at the expected size of 1.05 kb showing that the line 24C contained the HPT coding sequence. No band was observed for DNA from control tissue. The name of the callus line 24C was changed to PH2.

To demonstrate that the hygromycin gene is incorporated into high molecular weight DNA, DNA isolated from PH2 callus and control callus was treated with (i) no restriction enzyme, (ii) BamHI, as described previously, or, (iii) PstI, which cuts the plasmid 35 pHYGI1 only once within the HPT coding sequence. Samples were blotted and probed with the HPT coding sequence as described previously.

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Undigested PH2 DNA only showed hybridization to the probe at the position of uncut DNA, demonstrating that the hygromycin gene is incorporated into high molecular weight DNA. The expected 1.05 kb band for PH2 DNA digested with BamHI was observed, as had been shown previously. For PH2 DNA digested with PstI, a 5.9 kb band would be expected if the hygromycin gene was present on an intact pHYGI1 plasmid. Two or more bands of variable size (size dependent on the position of 10 flanking PstI sites within the host DNA) would be expected if the gene was integrated into genomic DNA. Two bands were observed with approximate molecular sizes of 6.0 and 3.0 kb. These data are consistent with incorporation of the hygromycin gene into high molecular 15 weight DNA. No hybridization was observed for DNA from control callus in any of the above treatments.

These results demonstrate that the HPT coding sequence is not present in PH2 callus as intact pHYGI1 or as a small non-chromosomal plasmid. They are consistent with incorporation of the hygromycin gene into high molecular weight DNA. Further Southern blot analyses demonstrated that the HPT coding sequence is contiguous with the 35S promoter sequence.

The line PH2, along with unselected control callus, was placed onto RM5 medium to regenerate plants as in Example I. After 16 days, the callus was transferred to R5 medium as in Example I. After 25 d on R5 medium, plantlets were transferred to R5 medium and grown up for 20 days. At this point, plantlets were transferred to vermiculite for one week and then transplanted into soil where they were grown to sexual maturity. Controlled pollinations were then performed

with the inbred line B73.

VII. Analysis of R1 progeny

A. PH2 as Pollen Donor

Ten R1 progeny plants from a B73 X PH 2.6 cross were assayed for HPT expression with both root

5 elongation and etiolated leaf assays. One positive plant was identified. The presence of the HPT gene in the positive plant was confirmed by Southern blot analysis employing the HPT probe as described in Example I.

To conduct the root elongation bioassay, seed was sterilized in a 1:1 dilution of commercial bleach in water plus 0.1% Alconox for 20 min. in 125 ml Erlenmeyer flasks and rinsed 3 times in sterile water. The seeds were imbibed overnight in sterile water containing 50 mg/ml Captan by shaking at 150 rpm.

After imbibition, the solution was decanted from the flasks and the seed transferred to flow boxes (Flow Laboratories) containing 3 sheets of H₂O saturated germination paper. A fourth sheet of water saturated 20 germination paper was placed on top of the seed. Seed was allowed to germinate 4 days.

After the seed had germinated, approximately 1 cm of the primary root tip was excised from each seedling and plated on medium containing MS salts, 20 g/l sucrose, 50 mg/l hygromycin, 0.25% Gelrite, and incubated in the dark at 26°C for 4 days.

Roots were evaluated for the presence or absence of abundant root hairs and root branches. Roots were classified as transgenic (hygromycin resistant) if they had root hairs and root branches, and untransformed (hygromycin sensitive) if they had limited numbers of branches.

After the root tips were excised, the seedlings of one PH2 ear and one control ear were transferred to 35 moist vermiculite and grown in the dark for 5 days. At this point, to conduct an etiolated leaf bioassay, 1 mm sections were cut from the tip of the coleoptile,

surface sterilized 10 seconds, and plated on MS basal salts, 20 g/l sucrose, 2.5 g/l Gelrite with either 0 (control) or 100 mg/l hygromycin and incubated in the dark at 26°C for 18 hrs. Each plate contained duplicate sections of each shoot. The plates were then incubated in a light regimen of 14 hrs light 10 hrs dark at 26°C for 48 hrs, and rated on a scale of from 0 (all brown) to 6 (all green) for the percent of green color in the leaf tissue. Shoots were classified as untransformed (hygromycin sensitive) if they had a rating of zero and classified as transformed (hygromycin resistant) if they had a rating of 3 or greater.

B. PH2 as Pollen Recipient

15 80 R1 progeny plants from a PH2 plant designated PH 2.23 X B73 cross were assayed for HPT expression by root elongation assay and 26 tested positive. The expression of the HPT gene was further confirmed for all 26 by direct exposure of seedlings to 20 hygromycin-containing medium in Flow boxes. Two of the resistant plants were analyzed by PCR, and the presence of the HPT sequence was confirmed.

VIII. Analysis of R2 Progeny

25 The transgenic R1 plant which had PH2 as the pollen parent was grown to maturity and used to pollinate an FR4326 plant.

R2 seeds from this cross were germinated and tested for the presence of HPT sequences by PCR.

30 Expression of the HPT gene was determined with the HPT enzymatic assay described in Example I.

Three of the 19 progeny tested contained HPT gene sequences and also expressed the HPT enzyme activity. The remainder neither contained HPT sequences nor showed enzyme activity. This result demonstrates pollen transmission and expression of the genetically engineered DNA through 2 successive generations,

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indicating that the gene is stable.

EXAMPLE V

The friable, embryogenic maize callus culture 5 AB12 described in Example III was used.

The plasmids pCHN1-1 and pLUC-1 were constructed in the vector pBS+ (Stratagene, Inc., San Diego, CA), a 3.2 Kb circular plasmid, using standard recombinant DNA techniques. pCHN1-1 contains the hygromycin B phosphotransferase (HPT) coding sequence from E. coli (Gritz et al. 1983) flanked at the 3' end by the nopaline synthase (nos) polyadenylation sequence of Agrobacterium tumefaciens (Chilton and Barnes 1983). Expression is driven by the cauliflower mosaic virus (CaMV) 35S promoter (Guilley et al. 1982), located upstream from the HPT coding sequence. The plasmids pBII221 and pHYGI1 were also used in this example.

pLUC-1 contains the firefly luciferase coding sequence (DeWet et al. 1987) flanked at the 5' end by the CaMV 35S promoter and at the 3' end by the nos polyadenylation sequence. This plasmid was used solely as negative control DNA. Plasmids were introduced into AB12 by microprojectile bombardment. Twenty-six plates were prepared as described in Example I. One tube contained only plasmid pBII221; two tubes contained both plasmids pHYGI1 and pBII221; two tubes contained both plasmids pCHN1-1 and pBII221; and one tube contained only plasmid pLUC-1. The bombardments which were performed are summarized on Table 7.

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Table 7

5	2 x pBII221 prep	To determine transient expression frequency
10	10 x pHYGI1/pBII221	As a potential positive treatment for transformation
	10 x pCHN1-1/pBII221	As a potential positive treatment for transformation
15	4 x pLUC-1	Negative control treatment

The two plates of callus bombarded with pBII221 20 were assayed for transient GUS expression. The number of blue cells was counted, giving 291 and 477 transient GUS expressing cells in the two plates, suggesting that the DNA delivery process had also occurred with the other bombarded plates.

Immediately after all samples had been bombarded, callus from all of the plates treated with pHYGI1/pBII221, pCHN1-1/pBII221 and three of the plates treated with pLUC-1 were transferred plate for plate onto round 1 selection plates, F medium containing 15 30 mg/l hygromycin B, (five pieces of callus per plate). Callus from the fourth plate treated with pLUC-1 was transferred to F medium without hygromycin. This tissue was subcultured every 2-3 weeks onto nonselective medium and is referred to as unselected control callus.

After two weeks of selection, tissue appeared essentially identical on both selective and nonselective media. All callus from eight plates from each of the pHYGI1/pBII221 and pCHN1-1/pBII221 treatments and two plates of the control callus on selective media were 40 transferred from round 1 selection plates to round 2 selection plates that contained 60 mg/l hygromycin. round 2 selection plates each contained ten 30 mg pieces

of callus per plate, resulting in an expansion of the total number of plates.

The remaining tissue on selective media, two plates each of pHYGI1/pBII221 and pCHN1-1/pBII221

5 treated tissue and one of control callus, were placed in GUS assay buffer at 37°C to determine whether blue clusters of cells were observable at two weeks postbombardment. After 6 days in assay buffer, this tissue was scored for GUS expression. The results are summarized on Table 8.

	Table 8	
<u>Treatment</u>	Replicate	<u>Observations</u>
pLUC-1		No blue cells
pHYGI1/pBII221	Plate 1	11 single cells 1 four-cell cluster
	Plate 2	5 single cells
pCHN1-1/pBII221	Plate 1	<pre>1 single cell 2 two-cell clusters</pre>
	Plate 2	5 single cells 1 two-cell cluster 2 clusters of 8-10 cells

After 21 days on the round 2 selection plates, all viable portions of the material were transferred to round 3 selection plates containing 60 mg/l hygromycin. The round 2 selection plates, containing only tissue that was apparently dead, were reserved. Both round 2 and 3 selection plates were observed periodically for viable proliferating sectors.

After 35 days on round 3 selection plates, both the round 2 and round 3 sets of selection plates were checked for viable sectors of callus. Two such sectors were observed proliferating from a background of dead tissue on plates treated with pHYGI1/pBII221. The first

sector named 3AA was from the round 3 group of plates and the second sector named PH1 was from the round 2 group of plates. Both lines were then transferred to F medium without hygromycin.

After 19 days on F medium without hygromycin, the line 3AA grew very little whereas the line PH1 grew rapidly. Both were transferred again to F medium for 9 days. The lines 3AA and PH1 were then transferred to F medium containing 15 mg/l hygromycin for 14 days. At this point, line 3AA was observed to be of very poor quality and slow growing. The line PH1, however, grew rapidly on F medium with 15 mg/l hygromycin; the line was then subcultured to F medium without hygromycin.

After 10 days on F medium, an inhibition study
of the line PH1 was initiated. Callus of PH1 was
transferred onto F medium containing 1, 10, 30, 100, and
250 mg/l hygromycin B. Five plates of callus were
prepared for each concentration and each plate contained
ten approximately 50 mg pieces of callus. One plate of
unselected control tissue was prepared for each
concentration of hygromycin.

It was found that the line PH1 was capable of sustained growth over 9 subcultures on 0, 10, 30, 100, and 250 mg/l hygromycin.

Additional sectors were recovered at various time points from the round 2 and 3 selection plates.

None of these were able to grow in the presence of hygromycin for multiple rounds, i.e., two or three subcultures.

To show that the PH1 callus had acquired the hygromycin resistance gene, a Southern blot of PH1 callus was prepared by isolating DNA from PH1 and unselected control calli by freezing 2 g of callus in liquid nitrogen and grinding it to a fine powder which was transferred to a 30 ml Oak Ridge tube containing 6 ml extraction buffer (7M urea, 250 mM NaCl, 50 mM TrisHCl pH 8.0, 20 mM EDTA pH 8.0, 1% sarcosine). To this

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was added 7 ml of phenol:chloroform 1:1, the tubes shaken and incubated at 37°C for 15 min. Samples were centrifuged at 8K for 10 min. at 4°C. The supernatant was pipetted through miracloth (Calbiochem 475855) into 5 a disposable 15 ml tube (American Scientific Products, C3920-15A) containing 1 ml 4.4 M ammonium acetate, pH Isopropanol, 6 ml was added, the tubes shaken, and the samples incubated at -20°C for 15 min. The DNA was pelleted in a Beckman TJ-6 centrifuge at the maximum 10 speed for 5 min. at 4°C. The supernatant was discarded and the pellet was dissolved in 500 μ l TE-10 (10 mM Tris-HCl pH 8.0, 10 mM EDTA pH 8.0) 15 min. at room temperature. The samples were transferred to a 1.5 ml Eppendorf tube and 100 μ l 4.4 M ammonium acetate, pH 5.2 15 and 700 μ l isopropanol were added. This was incubated at -20°C for 15 min. and the DNA pelleted 5 min. in an Eppendorf microcentrifuge (12,000 rpm). The pellet was washed with 70% ethanol, dried, and resuspended in TE-1 (10 mM Tris-HCl pH 8.0, 1 mM EDTA).

Ten μ g of isolated DNA were digested with BamHI (NEB) and analyzed as described in Example I using the HPT probe. As shown in Figure 3, a band was observed for PH1 callus at the expected position of 1.05 Kb, indicating that the HPT coding sequence was present. No 25 band was observed for control callus.

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To demonstrate that the hygromycin gene is incorporated into high molecular weight DNA, DNA isolated from PH1 callus and control callus was treated with (i) no restriction enzyme, (ii) BamHI, as described 30 previously, or (iii) PstI, which cuts the plasmid pHYGI1 only once within the HPT coding sequence. Samples were blotted and probed with the HPT coding sequence as described previously.

Undigested PH1 DNA only showed hybridization to 35 the probe at the position of uncut DNA, demonstrating that the hygromycin gene is incorporated into high molecular weight DNA. The expected 1.05 Kb band for PH1 DNA digested with BamHI was observed, as had been shown previously. For PH1 DNA digested with PstI, a 5.9 Kb band would be expected if the hygromycin gene was present on an intact pHYGI1 plasmid. Two or more bands of variable size (size dependent on the position flanking PstI sites within the host DNA) would be expected if the gene was incorporated into high molecular weight DNA. Three bands were observed with approximate molecular sizes of 12, 5.1, and 4.9 Kb.

- This result demonstrates incorporation of the hygromycin gene into high molecular weight DNA. The intensity of the 4.9 Kb band is approximately twice as great as the other two bands, suggesting either partial digestion or possibly a tandem repeat of the HPT gene. No
- 15 hybridization was observed for DNA from control callus in any of the above treatments.

These results demonstrate that the HPT coding sequence is not present in PH1 callus as intact pHYGI1 or as a small non-chromosomal plasmid. They are consistent with incorporation of the hygromycin gene into high molecular weight DNA. Further, Southern blot analyses demonstrated that the HPT coding sequence is contiguous with the 35S promoter sequence.

PH1 callus was transferred from all of the concentrations of hygromycin used in the inhibition study
to RM5 medium and plants were regenerated as described
in Example I. A total of 65 plants were produced from
PH1 and a total of 30 plants were produced from control
callus.

To demonstrate that the introduced DNA had been retained in the RO tissue, a Southern blot was performed as previously described on BamHI digested leaf DNA from three randomly chosen RO plants of PHI. The blot was probed with the HPT probe as in Example I. As shown in Figure 4, a 1.05 Kb band was observed with all three plants indicating that the HPT coding sequence was present. No band was observed for DNA from a control

plant.

Controlled pollinations of mature PH1 plants were conducted by standard techniques with inbred Zea mays lines A188, B73, and Oh43. Seed was harvested 45 days post-pollination and allowed to dry further 1-2 weeks.

The presence of the hygromycin resistance trait in the R1 progeny was evaluated by the root elongation bioassay, an etiolated leaf bioassay, and by Southern 10 blotting. Two ears each from regenerated PH1 and control plants were selected for analysis. The pollen donor was inbred line A188 for all ears. The results are shown in Fig. 5 and in Table 9, below.

Table 9

ANALYSIS OF PH1 R1 PLANTS

PH1 <u>PLANT</u>	ROOT ASSAY	LEAF ASSAY	BLOT	CONT PLANT	ROOT ASSAY	LEAF <u>ASSAY</u>	BLOT
3.1	+	ND	+	4.1	-	ND	ND
3.2	-	ND	-	4.2	-	ND	ND
3.3	-	ND	-	4.3	-	ND	ND
3.4	-	ND	-	4.4	-	ND	ND
3.5	-	ND	-	4.5	-	ND	ND
3.6	+	ND	+	4.6	-	ND	ND
3.7	-	ND	_	4.7	-	ND	ND
				2.1	-	ND	-
10.1	+	+	+	1.1	-	-	-
10.2	+	+	+	1.2	-	-	ND
10.3	-	-	ND	1.3	- '	-	ND
10.4	-	-	-	1.4	-	-	ND
10.5	-	-	-	1.5	-	_	ND
10.6	-	-	-	1.6	-	_	ND
10.7	-	-	-	1.7	-	_	ND
10.8	ND	+	+	1.8	_	_	ND

Key: + = transgenic; - = nontransgenic; ND = not done

The presence of the HPT gene was confirmed in two R2 progeny deriving from a PH1 maternal parent.

Plant PH1.3.1 (see Table 9) was pollinated with Oh43 pollen. Nine seeds derived from this cross were germinated, DNA was prepared from the leaves of four of the progeny plants, and analyzed by Southern blotting using the HPT coding sequence probe. Two of the four plants tested contained the HPT sequence in high copy number. Although the HPT gene was present in two of the plants, expression of the gene could not be detected in the etiolated leaf assay.

Plant PH1.10.8 (Table 9) was used to pollinate B73 and was also selfed. Fifty of the progeny from the outcross were tested for HPT expression in both root and leaf assays. No expression was detected. Likewise Southern blots on DNA from eight of the progeny did not detect the presence of the HPT gene. In the progeny from the self cross, no evidence for the presence of the gene was obtained in leaf assays of nine progeny, or from Southern blots of the DNA from four of these plants.

The recombinant DNA was only shown to be inherited by progeny when a transgenic plant (both R0 and R1) was used as a female. This is suggestive of maternal inheritance of the recombinant DNA for PH1.

All of the publications and patent documents cited hereinabove are incorporated by reference herein. The invention has been described with reference to various specific and preferred embodiments and techniques. However, it should be understood that many variations and modifications may be made while remaining within the spirit and scope of the invention.

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International Application No: PCT/ /
MICROORGANISMS
Optional Sheet in connection with the microorganism referred to on page, line of the description 1
A. IDENTIFICATION OF DEPOSIT 1
Further deposits are identified on an additional sheet : Plasmid, pHYGII; Plasmid, PZ27Z10
Name of depositary institution 4
AMERICAN TYPE CULTURE COLLECTION
Address of depositary institution (including postal code and country) • 12301 Parklawn Drive Rockville, Maryland 20852 U.S.A.
Date of deposit * Accession Number *
March 16, 1990 and December 20, 1990 40774 and 40938, respectively
B. ADDITIONAL INDICATIONS 7 (leave blank if not applicable). This information is continued on a separate attached sheet
C. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE 2 (If the indications are not for all designated States)
D. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)
The indications listed below will be submitted to the international Bureau later * (Specify the general nature of the indications e.g.,
"Accession Number of Deposit")
E. This enset was received with the international application when filed to be checked by the receiving Office)
(Authorized Officer) The date of receipt (from the applicant) by the International Bureau 14
was (Authorized Officer)

Form PCT RO-134 (January 1981)

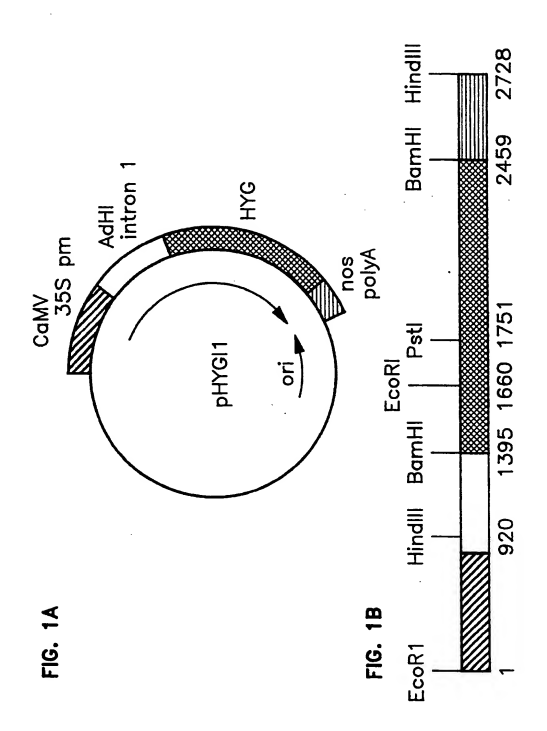
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WHAT IS CLAIMED IS:

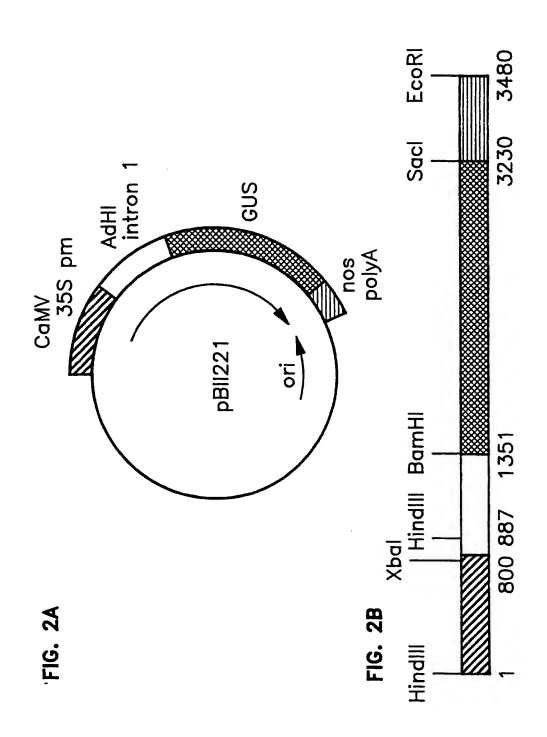
- 1. A fertile transgenic <u>Zea mays</u> plant containing recombinant DNA which is heritable.
- 2. The plant of claim 1 wherein the recombinant DNA is chromosomally integrated.
- 3. The plant of claim 1 wherein the recombinant DNA comprises a promoter.
- 4. The plant of claim 1 wherein the recombinant DNA is expressed.
- 5. The plant of claim 1 wherein the recombinant DNA encodes a protein.
- 6. The plant of claim 5 containing an effective amino acid-elevating amount of an expressed recombinant gene encoding a seed storage protein, which gene is heritable.
- 7. The plant of claim 6 wherein the recombinant gene encodes a 10 kD zein protein, which is expressed so that the level of whole kernel methionine is substantially elevated above the whole kernel methionine level in the corresponding Zea mays plant wherein said recombinant gene is absent.
- 8. The plant of claim 1 wherein said recombinant gene expresses a selectable marker gene or a reporter gene.
- 9. The plant of claim 8 wherein the selectable marker gene confers resistance or tolerance to a herbicide.

- 10. The plant of claim 8 wherein the selectable marker gene confers resistance or tolerance to a compound selected from the group consisting of hygromycin, kanamycin, G418, 2,2-dichloropropionic acid, neomycin, sethoxydim, haloxyfop, phosphinothricin, glyphosate, methotrexate, an imidazolinone herbicide, a sulfonylurea herbicide, a triazine herbicide and bromoxynil.
- 11. The plant of claim 10 wherein the selectable marker gene confers resistance or tolerance to hygromycin.
- 12. The plant of claim 8 which expresses a reporter gene.
- 13. The plant of claim 1 wherein said recombinant DNA comprises a DNA sequence selected from the group consisting of a bacterial dap A gene, a Bt-endotoxin gene, a protease inhibitor gene, a bacterial ESPS synthase gene, a chitinase gene, a glucan endo-1,3-β-glycosidase gene, structural DNA, a regulatory sequence, an identification sequence and a sequence encoding antisense RNA.
- 14. The seed produced by the plant of claims 1, 6, 8 or 13 which has inherited the recombinant gene.
- 15. The R1 and subsequent generations derived from the plant of claims 1, 6, 8 or 13.
- 16. The transgenic <u>Zea mays</u> plant of claim 1 containing recombinant DNA which is heritable, wherein said DNA was introduced into said plant, or an ancestor of said plant, at an embryonic stage.

- 17. The transgenic Zea mays plant of claim 16 wherein the recombinant DNA was introduced into said plant by microprojectile bombardment.
- 18. The transgenic Zea mays plant of claim 17 wherein the recombinant DNA was introduced by microprojectile bombardment of a callus culture.
- 19. A process for producing a fertile transgenic Zea mays plant which stably expresses recombinant DNA which is heritable, wherein the process comprises the steps of:
 - (i) establishing a regenerable cell culture from a Zea mays plant to be transformed;
 - (ii) transforming cells from said culture by bombarding said cells with recombinant DNA-coated microprojectiles;
 - (iii) identifying or selecting transformed
 cells; and
 - (iv) regenerating a fertile transgenic <u>Zea mays</u> plant therefrom, which stably expresses said recombinant DNA which is heritable.
- 20. The process of claim 19 wherein the cell culture is a friable, embryonic callus culture.
- 21. The process of claim 20 wherein the callus culture subjected to bombardment is in clumps of about 30 to 80 mg per clump.
- 22. The process of claim 21 wherein said callus is initiated on solid media.
- 23. A plant prepared by the process of claim 19 wherein the plant is regenerated from transformed callus.



SUBSTITUTE SHEET



SUBSTITUTE SHEET

FIG. 3

PHI CALLUS

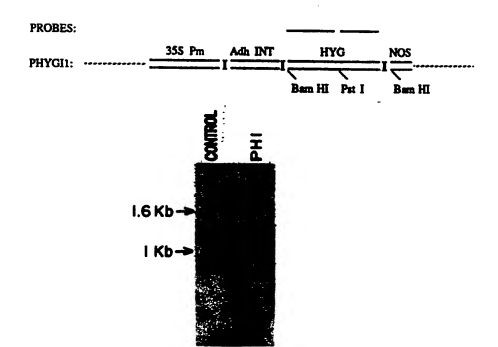


FIG. 4

PHI RO PLANTS

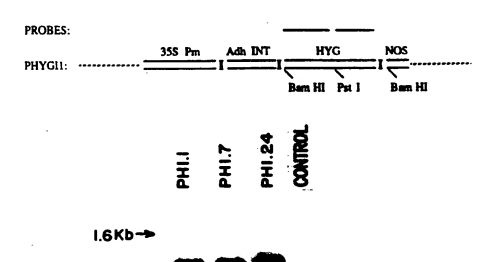


FIG. 5

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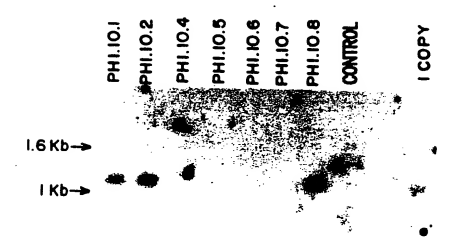
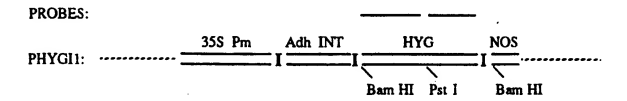
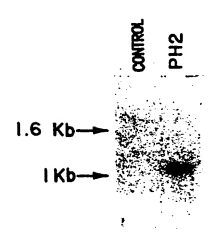
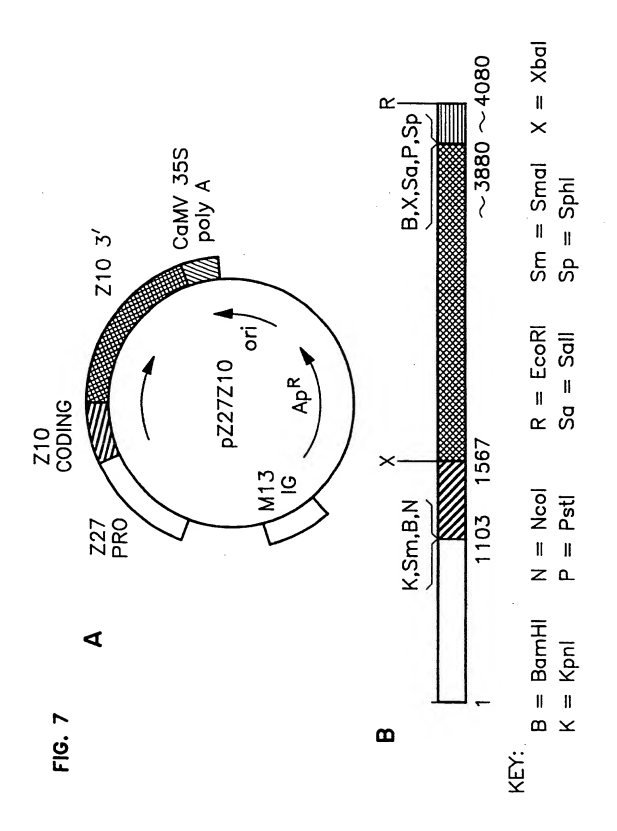


FIG. 6 PH2 CALLUS





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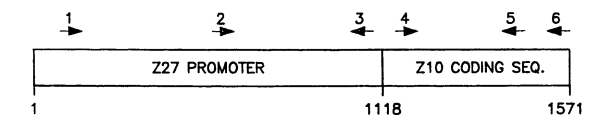


FIG. 8

INTERNATIONAL SEARCH REPORT

International Application 40. PCT/US91/00183

I. CLASSI	FICATIO	N OF SUBJECT MATTER (if several class	Michiga symbols apply underly all 6	
According t	o internati	onal Patent Classification (IPC) or to both Na	itional Classification and ISC	
IPC(5	5):C12	N 500, 15/00; 15/11; AO1H	4.700	
U.S.C	L : 4	35/240.4, 172.3, 320; 536	/27: 800/205	
II. FIELDS	SEARCH	IED	, 51, 600, 203	
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		435/240.4, 172.3, 3	20; 536/27, 800/205	
U.S.				
	•	Documentation Searched other	than Minimum Documentation	
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III. DOCUM	ENTS C	ONSIDERED TO BE RELEVANT 1		
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IV. CERTIFI			and the state of the study to	There's
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. OB	SERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1	
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	national search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons; m numbers because they relate to subject matter not required to be searched by this Authority, namely;	
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his Inter	national Searching Authority found multiple inventions in this international application as follows:	
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	all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims he international application.	
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cla	aim 11 (hygromycin)and claim 13 (bacterial dep A gene). Telephone pract	ice
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As Invi	all searchable claims could be searched without effort justifying an additional fee, the international Searching Authority did not to payment of any additional fee.	
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OBSERVATION WHERE UNITY OF INVENTION IS LACKING

Detailed Reasons for Holding Lack of Unity of Invention

In the examination of international applications filed under the Patent Cooperation Treaty, PCT Rule 13.1 states that the international application shall relate to one invention only or to a group of inventions so linked as to form "a single general inventive concept."

PCT Rule 13.2 indicates that this shall be construed as permitting, in particular, one of the following three possible combinations of claimed invention:

- (1) a product, a process specifically adapted for the manufacture of said product and a use of said product, or
- (2) a process, and an apparatus or means specifically designed for carrying out said process, or
- (3 a product, a process specially adapted for the manufacture of said product and an apparatus or means designed for carrying out the process.

Additionally, current United States Patent and Trademark Office restriction practice permits the following combinations of claimed invention:

- (4) a product, and a process specifically adapted for the manufacture of said product, and
- (5) a product, and a use of said product, as where said use ass claimed cannot be practiced with another materially different product.

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept. In order for all inventions to be

examined, the appropriate additional examination fees must be paid.

The claims present a plurality of multiply distinct inventions as follows:

I. Claims 1-2 and 15-18 are drawn to the product of a fertile transgenic Zea mays plant and are, for example, classified in Class 800, subclasses 205. In addition, claims 19-23 are drawn to the process of making the above product and are, for example, classified in Class 435, subclass 240.4.

Note that the following are independent and distinct species pertinent to the inventions of Group I where a) is the first species and will be searched with claims 1-2, 15-18 and 19-23 in the event that no other fees are paid. Note that the search of any other additional species with Group I requires payment of additional fees.

- a) the transgenic plant wherein the recombinant DNA comprises a promoter (claim 3).
- b) the transgenic plant wherein the recombinant DNA is expressed (claim 4).
- c) the transgenic plant wherein the recombinant gene encodes a seed storage protein (claims 5-7).
- d) the transgenic plant wherein the recombinant gene expresses a selectable marker (claims 8-11) from the group consisting of:
 - d1) hygromycin, kanamycin, G418, 2,2-dichloropropionic acid, neomycin, sethoxydim, haloxyfop, phosphinothricin,

glyphosate, methotrexate, an imidazolinone herbicide, a sulfonylurea herbicide, a triazolopyrimidine herbicide, a triazine herbicide or bromoxynil.

If this species is elected a further election of one of the following subspecies is required. If no subspecies is elected the first subspecies will be automatically searched. In addition, as claim 11 is limited to hygromycin election of a subspecies other than hygromycin would require a further payment for an additional group.

- e) the transgenic plant wherein the recombinant gene expresses a reporter gene (claims 8 and 12).
- f) the transgenic plant wherein the recombinant DNA comprises a gene sequence selected from the group (claim 13):
 - f1) bacterial <u>dap</u> A, Bt-endotoxin, protease inhibitor, bacterial ESPS synthetase, chitinase, glucan endo-1-3-B-glycosidase, a structural DNA, regulatory sequence, identification sequence or a sequence encoding antisense RNA.
- II. Claim 14 is drawn to seed produced by the plants of claims 1, 6, 8 or 13 and are for example, classified in Class 800, subclasses 200.

Note that the following are independent and distinct species pertinent to the inventions of Group II where a) is the first species and will be searched with claim 14 in the event that no other fees are paid. Note that the search of any other additional species with Group I requires payment of additional fees.

- a) seed of the fertile transgenic <u>Zea mays</u> containing heritable recombinant DNA (claim 1).
- b) seed of the fertile transgenic <u>Zea mays</u> containing an effective amino acid-elevating amount of an expressed seed storage protein (claim 6).
- c) seed of the fertile transgenic <u>Zea mays</u> containing a recombinant selectable gene (claim 8).
- d) seed of the fertile transgenic <u>Zea mays</u> containing a recombinant reporter gene (claim 8).
- e) seed of the fertile transgenic <u>Zea mays</u> containing a recombinant DNA sequence selected from the group consisting of:
- e1) bacterial <u>dap</u> A, Bt-endotoxin, protease inhibitor, bacterial ESPS synthetase, chitinase, glucan endo-1-3-B-glycosidase, a structural DNA, regulatory sequence, identification sequence or a sequence encoding antisense RNA (claim 13).

The inventions listed as Groups I and II do not meet the requirements for Unity of Invention for the following reasons:

1) the product of Group I — the transgenic plant — and its method of making is separately usable and consists of a distinctly different invention — the seed of the transgenic plant — of Group II. Given the plant and its method of making as defined by the claims the effect and use of transgenic seeds would not necessarily be obvious.

Regarding the requirement for the additional election of a species as defined in Group I, subspecies d1 and e1 and Group II, subspecies e1; the inventions as defined by each subspecies are deemed distinct since there is nothing on this record to show them to be obvious variants. Should applicant traverse on the ground that the species are not distinct, applicant should submit evidence or identify such evidence now of record showing the species to be obvious variants or clearly admit on the record that this is the case. Databases: Dialog Information Services On-line (File set AGRI) STN International (Files set Biosis and Chem Abstracts), Automated Patent System (File USPAT)

PCT/US91/00183
Attachment to Form PCT/ISA/210

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II. Search terms: Transgenic with maize or corn or Zea mays; Gene expression with zein storage protein, seed storage protein, reporter gene, selectable marker gene, herbicide resistance; Transformation and microprojectile or biolistic; and callus culture or in vitro; hygromycin or bacterial dap A gene.

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III. DOCUMENTS CONSIDERED TO BE RELEVANT

<u>Cate</u>	gory Citation	Claims
Y	Nature, Volume 327, issued07 May 1987, (London GB) Klein et al. "High-velocity microprojectiles for delivering nucleic acids into living cells" pages 69-73, see the entire article.	17-23
Y	Proc. Nat. Acad. Sci., Volume 85 issued June 1988 (Washington DC) Klein et al. "Transfer of foreign genes into maize cells with high-velocity microprojectiles" pages 4305-4309, see the entire article.	17-23
Y	Bio/technology, Volume 6, issued May 1988, (New York NY) Klein et al., "Factors influencing gene delivery into Zea mays cells by high-velocity microprojectiles" pages 559-563 in particular the Results and Discussion at pages 559-562.	17-23
· · · • Y	Plant Physiol., Volume 91, issued 1989, (Rockville MD) Klein et al. "Genetic transformation of maize cells by particle bombardment" pages 440-444, see the entire article.	17-23
Y	Bio/technology, Volume 6, issued January 1988, (New York NY) Rhodes et al. "Plant regeneration from protoplasts isolated from embryogenic maize cell cultures" pages 56-60 in particular the Discussion at page 59.	1-23
. ү	Science, Volume 240, issued 08 April 1988, (Washington DC) Rhodes et al. "Genetically transformed plants from protoplasts" pages 204-207, see the entire article.	1-5, 10, 12
А	Particulate Science and Technology, Volume 5 issued 1987, Sanford et al. "Delivery of substances into Cells and tissue using a particle bombardment process" pages 27-37 in particular page 35.	17-23

Y	Plant Physiol., Volume 86 issued 1988 (Rockville MD) Hauptmann et al. "Evaluation of selectable Markers for obtaining stable transformants in the Gramineae" pages 602-606 in particular the Discussion at pages 604-605.	8-11
A	Trends in Biotechnology, Volume 6 issued December 1988, (Amsterdam Netherlands) Sanford, J. "The biolistic process" pages 299-302, see the entire article.	17-23
Y	Bio/technology, Volume 7 issued June 1989, (New York NY) Shillito et al. "Regeneration of fertile plants from protoplasts of elite inbred maize" pages 581-587 in particular the Discussion at page 584-585.	1-23
Y	Plant Molecular Biology, Volume 7 issued 1986, (Dordrecht, NL) Graves et al. "The transformation of <u>Zea mays</u> seedlings with <u>Agrobacterium tume</u> -faciens" pages 43-50, see entire document.	1-5
Y	Bio/technology, Volume 7, issued June 1989, (New York NY) Prioli et al. "Plant regeneration and recovery of fertile plants from protoplasts of maize (Zea mays L.)" pages 589-594 in particular the Discussion at pages 589 and 594.	1-16
Υ, Ο	J. Cellular Biochem, 27 March - 7 April 1989, Gordon-Kamm et al. "Stable transformation of Embryogenic maize cultures by microprojectile bombardment" Suppl 13D, page 305 Abstract M122.	1-23
Υ, Ο	J. Cellular Biochem, 27 March - 7 April 1989, Ross et al. "Transient and stable transgenic cells and calli of tobacco and maize following microprojectile bombardment" Suppl 13D, page 268 Abstract M149.	1-23
$\frac{X}{Y}$	EF,A,D242,435 (Rice et al.) 23 November 1986, see entire document.	<u>1-5</u> 6-18
Υ .	Nature, Volume 338, issued 16 March 1989, (London GB) Shimamoto et al. "Fertile transgenic rice plants regenerated from transformed protoplasts" pages 274-276, see the entire article.	8-12

<u>X</u> Y	Jap. J. Breed., Volume 30, Suppl. 2, issued 1980, Ohta et al. "Gene manifestation of exogenous DNA applied to self-propagating stigma" pages 184-185 (translation pages 1-3) see entire article.	<u>1-5</u> 6-16
Y	Proc. Nat. Acad. Sci., Volume 83, issued February 1986, (Washington DC) Ohta et al. "High-efficiency genetic transformation of maize by a mixture of pollen and exogenous DNA" pages 715-719, see entire article.	1-16
Υ	Biological Abstracts, Volume 82, No 1 issued 01 July 1986, François et al. "Chromosomal location and nucleotide sequence of the Echerichia coli dapA gene" see column 1 abstract no. 3396 J. Bacteriol 166(1): 297-300 1986.	13
P,Ÿ	Trends in Biotech., Volume 8, issued 1990. (Amsterdam Netherlands) Christou et al. "Soybean genetic engineering - commercial production of transgenic plants, pages 145-151.	19-23
P,Y	Bio/technology, Volume 8, issued September 1990, (New York NY) Fromm et al. "Inheritance and expression of chimeric genes in the progeny of transgenic maize plants" pages 833-839.	1-23
P,Y	The Plant Cell, Volume 2 issued July 1990, Gordon-Kamm et al. "Transformation of maize cells and regeneration of fertile transgenic	1-23

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